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(54) Title: NOVEL CYTOKINE

(57) Abstract

There is disclosed a polypeptide (CD40-L) and DNA sequences, vectors and transformed host cells useful in providing CD40-L polypeptides. More particularly, this invention provides isolated human and murine CD40-L polypeptides that bind to the extracellular binding region of a CD40 receptor.

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TITLE

NOVEL CYTOKINE

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TECHNICAL FIELD OF THE INVENTION

The present invention relates to a novel cytokine. More specifically, the present invention relates to the cloning of a murine and a human cytokine that binds to a human CD40 having both agonist and antagonist activity in soluble and membrane-bound forms.

BACKGROUND OF THE INVENTION

Cytokines that have an "Interleukin" designation are those protein factors that influence immune effector cells. Cytokines designated interleukin-1 through interleukin-12 have been reported and named as an interleukin. Other known cytokines include tumor necrosis factor (TNF), granulocyte-macrophage colony stimulating factor (GM-CSF), granulocyte colony stimulating factor (G-CSF), mast cell growth factor (MGF), epidermal growth factor (EGF), platelet-derived growth factor (PDGF), nerve growth factor (NGF), erythropoietin (EPO), γ -interferon (γ -IFN) and others.

DNAs for two different TNF receptors (Type I and Type II) have been cloned (Smith et al., Science 248:1019, 1990; and Schall et al., Cell 61:361, 1990). Both forms of TNF receptor are related to each other and belong to a family of receptors whose members include nerve growth factor receptor (Johnson et al., Cell 47:545, 1986), B cell antigen CD40 (Stamenkovic et al., EMBO J. 8:1403, 1989), T cell antigen OX40 (Mallett et al., EMBO J. 9:1063, 1990), human Fas antigen (Itoh et al., Cell 66:233, 1991) and murine 4-1BB receptor (Kwon et al., Cell. Immunol. 121:414, 1989 [Kwon et al. I] and Kwon et al., Proc. Natl. Acad. Sci. USA 86:1963, 1989 [Kwon et al. II]).

Human CD40 protein (CD40) is a peptide of 277 amino acids having a molecular weight of 30,600, and a 19 amino acid secretory signal peptide comprising predominantly hydrophobic amino acids (Stamenkovic et al.). The molecular weight (exclusive of glycosylation) of the mature human CD40 protein is 28,300. A cDNA encoding human CD40 was isolated from a cDNA library prepared from Burkitt lymphoma cell line Raji. The putative protein encoded by the CD40 cDNA contains a putative leader sequence, trans-membrane domain and a number of other features common to membrane-bound receptor proteins. CD40 has been found to be expressed on B lymphocytes, epithelial cells and some carcinoma cell lines.

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A monoclonal antibody (mAb) directed against CD40 has been shown to mediate various functional effects of human B cells. These effects include: (a) homotypic adhesions (Gordon et al., *J. Immunol.* 140:1425, 1988 [Gordon et al. I]); (b) increased cell size (Gordon et al. I and Valle et al., *Eur. J. Immunol.* 19:1463, 1989); (c) proliferation of B cells activated with anti-IgM, anti-CD20 mAb, phorbol ester alone (Clark et al., *Proc. Natl. Acad. Sci. USA* 83:4494, 1986; and Paulie et al., *J. Immunol.* 142:590, 1989), or phorbol ester combined with interleukin-4 (Gordon et al., *Eur. J. Immunol.* 17:1535, 1987 [Gordon et al. II]; and (d) production of IgE (Jabara et al., *J. Exp. Med.* 172:1861, 1990; Zhang et al., *J. Immunol.* 146:1836, 1991) and IgM (Gascan et al., *J. Immunol.* 147:8, 1991) from interleukin-4 (IL-4) stimulated T-depleted cultures.

One such antibody, called mAb 89 by Banchereau et al., Clin. Immunol. Spectrum 3:8, 1991 [Banchereau et al. I], was found to induce human B cell proliferation at a relatively low antibody concentration (30 ng/ml or about 10⁻¹⁰ M). Proliferation lasted two to three weeks and resulted in a ten-fold expansion of the human B cell population. Optimal stimulation of the B cells occurred when CD40 surface molecule was cross-linked by IgM. Fab fragments of another anti-CD40 mAb induced only a weak proliferative response. Further, Banchereau et al., Science 251:70, 1991 [Banchereau et al. II] reported that resting human B cells entered a state of sustained proliferation when incubated with both a murine fibroblastic Ltk- cell line that was transfected with human Fc receptor and with a monoclonal antibody specific for human CD40. Banchereau et al. II found that cross-linking CD40 is necessary for clonal expansion of B cells.

CD23 is a low affinity IgE receptor that has been found to be expressed on most IgM-/IgD- mature B cells, but not T cells. CD23 has been sequenced and its sequence was described in Kikutani et al., Cell 47:657, 1986. Soluble CD23 (sCD23) was found to induce a pyrogenic reaction in rabbits and this reaction was abrogated by administration of human IgE (Ghaderi et al., Immunology 73:510, 1991). Therefore, CD23 may be an appropriate marker for soluble CD40 or CD40-L effects.

Prior to the present invention, a ligand for CD40 was unknown. Accordingly, there is a need in the art to identify and characterize a CD40 ligand (CD40-L).

SUMMARY OF THE INVENTION

A novel cytokine, hereafter referred to as "CD40-L," has been isolated and characterized. The nucleotide sequence and deduced amino acid sequence of representative murine CD40-L cDNA is disclosed in SEQ ID NO:1 and Figure 1, and the amino acid sequence is also listed in SEQ ID NO:2. The nucleotide sequence and

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deduced amino acid sequence of representative human CD40-L cDNA is disclosed in SEQ ID NO:11 and Figure 2, and the amino acid sequence is also listed in SEQ ID NO:12. The present invention further comprises other CD40-L polypeptides encoded by nucleotide sequences that hybridize, under moderate or severe stringency

5 conditions, to probes defined by SEQ ID NO:11 (the coding region of human CD40-L), fragments of the sequence extending from nucleotide 46 to nucleotide 828 of SEQ ID NO:11, or to DNA or RNA sequences complementary to Figure 2 (SEQ ID NO:11) or fragments thereof. The invention further comprises nucleic acid sequences which, due to the degeneracy of the genetic code, encode polypeptides substantially identical or substantially similar to polypeptides encoded by the nucleic acid sequences described above, and sequences complementary to them.

CD40-L is a type II membrane polypeptide having an extracellular region at its C-terminus, a transmembrane region and an intracellular region at its N-terminus. A soluble version of murine CD40-L has been found in supernatants from EL-4 cells and EL-4 cells sorted on the basis of a biotinylated CD40/Fc fusion protein described herein. Soluble CD40-L comprises an extracellular region of CD40-L or a fragment thereof. The protein sequence of murine CD40-L is described in Figure 1 and SEQ ID NO:2, and human CD40-L in Figure 2 and SEQ ID NO:12. The extracellular region of murine CD40-L extends from amino acid 47 to amino acid 260 in Figure 1 and SEQ ID NO:2, and of human CD40-L from amino acid 47 to amino acid 261 in Figure 2 and SEQ ID NO:12. CD40-L biological activity is mediated by binding of this cytokine with CD40 and includes B cell proliferation and induction of antibody secretion, including IgE secretion.

The present invention further provides antisense or sense oligonucleotides (deoxyribonucleotides or ribonucleotides) that correspond to a sequence of at least about 12 nucleotides selected from the nucleotide sequence of CD40-L or DNA or RNA sequences complementary to the nucleotide sequence of CD40-L as described in SEQ ID NO:1 and SEQ ID NO:11 and in Figures 1 and 2. Such antisense or sense oligonucleotides prevent transcription or translation of CD40-L mRNA or polypeptides.

Further still, the present invention provides CD40-L peptide fragments that correspond to a protein sequence of at least 10 amino acids selected from the amino acid sequence encoded by SEQ ID NO:1 or SEQ ID NO:11 that can act as immunogens to generate antibodies specific to the CD40-L immunogens. Such CD40-L immunogen fragments can serve as antigenic determinants in providing monoclonal antibodies specific for CD40-L.

The invention also provides a human CD40/Fc fusion protein and a soluble CD40 protein (sCD40) comprising the extracellular region of human CD40. Both

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sCD40 and CD40/Fc fusion protein can inhibit CD40-L or anti-CD40 mAb induced B cell stimulation, IL-4-induced IgE stimulation and IL-4 induced CD23 induction in B cells.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 illustrates nucleotide and amino acid sequences corresponding to murine CD40-L. This protein is a type II polypeptide having its N-terminus as its intracellular domain, followed by a transmembrane region, and an extracellular domain at the C-terminus of the polypeptide. The extracellular domain, which is longer than either the intracellular domain or the transmembrane region, contains one potential N-linked glycosylation site and two potential disulfide bonds in view of four cysteine (Cys) residues.

Figure 2 illustrates nucleotide and amino acid sequences corresponding to human CD40-L. This protein is a type II polypeptide having its N-terminus as its intracellular domain, followed by a transmembrane region, and an extracellular domain at the C-terminus of the polypeptide. The extracellular domain, which is longer than either the intracellular domain or the transmembrane region, contains 1 potential N-linked glycosylation site and 2 potential disulfide bonds in view of 5 cysteine (Cys) residues.

Figure 3 illustrates a comparison of protein sequences of human and murine CD40-L showing 77.7% homology at the amino acid level.

Figure 4 illustrates proliferation of T cell depleted human peripheral blood mononuclear cells (PBMC) caused by incubation with CV1 cells transfected with full length murine CD40-L cDNA (SEQ ID NO:1) and expressing bound CD40-L (CD40-L+CV1 cells) when compared with CV1 cells transfected with empty vector (HAVEO) and not expressing bound murine CD40-L. The day 7 proliferation results show that CD40-L+CV1 cells significantly increase proliferation of T- cell depleted PBMC in the presence or absence of interleukin-4 (IL-4).

Figure 5 illustrates a second determination of T cell depleted PBMC proliferation with addition of bound murine CD40-L and 10 ng/ml of IL-4. These data show no co-mitogenic effect of IL-4 but continued strong mitogenic effect of bound CD40-L.

Figure 6 illustrates that bound CD40-L augments IgE secretión.

Figure 7 illustrates that membrane-bound CD40-L stimulates CD23 shedding in the presence of IL-4.

Figure 8 illustrates proliferation of murine splenic B cells caused by membrane-bound murine CD40-L or 7A1 cells, which is a helper T cell clone.

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Figure 9 illustrates a comparison of murine EL40.9 cells, a sorted cell line that was sorted on the basis of expression of murine CD40-L and T cells 7A1 for induction of an antigen-specific response indicated by plaque forming cells (PFC) by anti-sheep red blood cells (SCBC).

Figure 10 illustrates a comparison of B cell proliferative activity of membrane-bound CD40-L and other cell types transfected with different cDNAs. Membrane-bound CD40-L showed significantly more B cell proliferative activity than a helper T cell clone or other control cells.

Figure 11 illustrates that 7C2 cells (a helper T cell clone) and CV1 cells transfected with murine CD40-L cDNA induce anti SRBC plaque forming cells.

Figure 12 illustrates a comparison of two helper T cell clones with cells expressing membrane-bound CD40-L for inducing murine B cell proliferation.

Figure 13 illustrates induction of antigen-specific plaque forming cells by membrane-bound CD40-L and a helper T cell clone in the presence or absence of added interleukin-2 (IL-2).

Figure 14 shows effects of membrane-bound CD40-L stimulating B cell proliferation and IgE secretion. The effects of membrane-bound CD40-L were inhibited by CD40 receptor but not by TNF receptor.

DETAILED DESCRIPTION OF THE INVENTION

Novel polypeptides that can act as a ligand for murine and human CD40 have been isolated and sequenced. More particularly, cDNAs encoding these ligands have been cloned and sequenced. Further provided are methods for expression of recombinant CD40-L polypeptides. CD40-L polypeptide include other forms of mammalian CD40-L, such as derivatives or analogs of human or murine CD40-L. Murine and human CD40-L comprise a 214 and 215, respectively amino acid extracellular region at the C-terminus of full length, membrane-bound polypeptide. The extracellular region contains the domain that binds to CD40. Murine and human CD40-L further comprise a homologous hydrophobic 24 amino acid transmembrane region delineated by charged amino acids on either side and a 22 amino acid intracellular region at their N-termini. The present invention further comprises full length CD40-L polypeptides or fragments thereof comprising all or part of the extracellular region or derivatives of the extracellular region and mammalian cells transfected with a cDNA encoding murine or human CD40-L and expressing human or murine CD40-L as a membrane-bound protein.

The present invention comprises isolated DNA sequences encoding CD40-L polypeptides and DNA or RNA sequences complementary to such isolated DNA

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sequences. The isolated DNA sequences and their complements are selected from the group consisting of (a) nucleotides 184 through 828, nucleotides 193 through 828 or nucleotides 193 through 762 of the DNA sequence set forth in Figure 2 (SEQ ID NO:11) and their complements, (b) DNA sequences which hybridize to the DNA sequences of (a) or their complements under conditions of moderate stringency and which encode a CD40-L polypeptide, analogs or derivatives thereof, and (c) DNA sequences which, due to the degeneracy of the genetic code, encode CD40-L polypeptides encoded by any of the foregoing DNA sequences and their complements. In addition, the present invention includes vectors comprising DNA sequences encoding CD40-L polypeptides and analogs, and host cells transfected with such vectors.

The novel cytokine disclosed herein is a ligand for CD40, a receptor that is a member of the TNF receptor super family. Therefore, CD40-L is likely to be responsible for transducing signal via CD40, which is known to be expressed, for example, by B lymphocytes. Full-length CD40-L is a membrane-bound polypeptide with an extracellular region at its C terminus, a transmembrane region, and an intracellular region at its N-terminus. A soluble version of CD40-L can be made from the extracellular region or a fragment thereof and a soluble CD40-L has been found in culture supernatants from cells that express a membrane-bound version of CD40-L. The protein sequence of the extracellular region of murine CD40-L extends from amino acid 47 to amino acid 260 in Figure 1 and SEQ ID NO:2. The protein sequence of the extracellular region of human CD40-L extends from amino acid 47 to amino acid 261 in Figure 2 and SEQ ID NO:12. The biological activity of CD40-L is mediated by binding to CD40 or a species-specific homolog thereof and comprises proliferation of B cells and induction of immunoglobulin secretion from activated B cells. CD40-L (including soluble monomeric and oligomeric forms, as well as membrane-bound forms) can effect B cell proliferation and immunoglobulin secretion (except IgE secretion) without the presence of added IL-4, in contrast to anti-CD40 antibodies, which require IL-4 and cross-linking to mediate activity.

CD40-L refers to a genus of polypeptides which are capable of binding CD40, or mammalian homologs of CD40. As used herein, the term "CD40-L" includes soluble CD40-L polypeptides lacking transmembrane and intracellular regions, mammalian homologs of human CD40-L, analogs of human or murine CD40-L or derivatives of human or murine CD40-L.

CD40-L may also be obtained by mutations of nucleotide sequences coding for a CD40-L polypeptide. A CD40-L analog, as referred to herein, is a polypeptide substantially homologous to a sequence of human or murine CD40-L but which has an

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amino acid sequence different from native sequence CD40-L (human or murine species) polypeptide because of one or a plurality of deletions, insertions or substitutions. Analogs of CD40-L can be synthesized from DNA constructs prepared by oligonucleotide synthesis and ligation or by site-specific mutagenesis techniques.

The primary amino acid structure of human or murine CD40-L may be modified to create CD40-L derivatives by forming covalent or aggregative conjugates with other chemical moieties, such as glycosyl groups, lipids, phosphate, acetyl groups and the like, or by creating amino acid sequence mutants. Covalent derivatives of CD40-L are prepared by linking particular functional groups to CD40-L amino acid side chains or at the N-terminus or C-terminus of a CD40-L polypeptide or the extracellular domain thereof. Other derivatives of CD40-L within the scope of this invention include covalent or aggregative conjugates of CD40-L or its fragments with other proteins or polypeptides, such as by synthesis in recombinant culture as N-terminal or C-terminal fusions. For example, the conjugate may comprise a signal or leader polypeptide sequence at the N-terminal region or C-terminal region of a CD40-L polypeptide which co-translationally or post-translationally directs transfer of the conjugate from its site of synthesis to a site inside or outside of the cell membrane or cell wall (e.g. the α-factor leader of Saccharomyces). CD40-L polypeptide fusions can comprise polypeptides added to facilitate purification and identification of CD40-L (e.g. poly-His), or fusions with other cytokines to provide novel polyfunctional entities. Other cytokines include. for example, any of interleukins-1 through 13, TNF (tumor necrosis factor), GM-CSF (granulocyte macrophage-colony stimulating factor), G-CSF (granulocyte-colony stimulating factor), MGF (mast cell growth factor), EGF (epidermal growth factor), PDGF (platelet-derived growth factor), NGF (nerve growth factor), EPO (erythropoietin), γ-IFN (gamma interferon), 4-1BB-L (4-1BB ligand) and other cytokines that affect immune cell growth, differentiation or function.

Nucleic acid sequences within the scope of the present invention include DNA and/or RNA sequences that hybridize to the nucleotide sequence of SEQ ID NO:1 or SEQ ID NO:11 or the complementary strands, under conditions of moderate or severe stringency. Moderate stringency hybridization conditions refer to conditions described in, for example, Sambrook et al. *Molecular Cloning: A Laboratory Manual*, 2 ed. Vol. 1, pp. 1.101-104, Cold Spring Harbor Laboratory Press, (1989). Conditions of moderate stringency, as defined by Sambrook et al., include use of a prewashing solution of 5 X SSC, 0.5% SDS, 1.0 mM EDTA (pH 8.0) and hybridization conditions of 50°C, 5 X SSC, overnight. Conditions of severe stringency include higher temperatures of hybridization and washing.

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Biological activity of CD40-L may be determined, for example, by competition for binding to the ligand binding domain of CD40 (i.e. competitive binding assays). Both murine CD40-L and human CD40-L bind to human CD40. The binding affinity of murine CD40-L (expressed on sorted murine EL-40.9 cells) for human CD40 was approximately

 $1.74 \times 10^9 \, \text{M}^{-1}$. Similarly, the binding affinity of murine CD40-L (expressed on unsorted murine EL-46.1 cells) for human CD40 was approximately $2.3 \times 10^9 \, \text{M}^{-1}$. Both binding affinity measurements are within a range typical of cytokine/cytokine receptor binding.

One configuration of a competitive binding assay for CD40-L polypeptide uses a radiolabeled, soluble murine CD40-L according to Figure 1 (SEQ ID NO:1) or human CD40-L according to Figure 2 (SEQ ID NO:11), and intact cells expressing CD40 (e.g., human B cells). Instead of intact cells, one could substitute soluble CD40 (such as a CD40/Fc fusion protein) bound to a solid phase through a Protein A or Protein G interaction with the Fc region of the fusion protein. A second configuration of a competitive binding assay utilizes radiolabeled soluble CD40 such as a CD40/Fc fusion protein, and intact cells expressing CD40-L. Alternatively, soluble CD40-L could be bound to a solid phase.

Competitive binding assays can be performed using standard methodology. For example, radiolabeled murine CD40-L can be used to compete with a putative CD40-L homolog to assay for binding activity against surface-bound CD40. Qualitative results can be obtained by competitive autoradiographic plate binding assays, or Scatchard plots may be utilized to generate quantitative results.

Competitive binding assays with intact cells expressing CD40 can be performed by two methods. In a first method, B cells are grown either in suspension or by adherence to tissue culture plates. Adherent cells can be removed by treatment with 5 mM EDTA treatment for ten minutes at 37° C. In a second method, transfected COS cells expressing membrane-bound CD40 can be used. COS cells or another mammalian cell can be transfected with human CD40 cDNA in an appropriate vector to express full length CD40 with an extracellular region exterior to the cell.

Alternatively, soluble CD40 can be bound to a solid phase such as a column chromatography matrix, or a tube or similar substrate suitable for analysis for the presence of a detectable moiety such as ¹²⁵I. Binding to a solid phase can be accomplished, for example, by obtaining a CD40/Fc fusion protein and binding it to a protein A or protein G surface.

Another means to measure the biological activity of CD40-L and homologs thereof is to utilize conjugated, soluble CD40 (for example, ¹²⁵I-CD40/Fc) in

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competition assays similar to those described above. In this case, however, intact cells expressing CD40-L, or soluble CD40-L bound to a solid substrate, are used to measure competition for binding of conjugated, soluble CD40 to CD40-L by a sample containing a putative CD40 homolog.

CD40-L may also be assayed by measuring biological activity in a B cell proliferation assay. Human B cells may be obtained from human tonsils by purification by negative selection and Percoll density sedimentation, as described by Defrance et al., *J. Immunol.* 139:1135, 1987. Burkitt lymphoma cell lines may be used to measure cell proliferation in response to CD40-L. Examples of Burkitt lymphoma cell lines include, for example, Raji (ATCC CCL 86), Daudi (ATCC CCL 213) and Namalwa (ATCC CRL 1432). Membrane-bound CD40-L stimulated B cell proliferation. Oligomeric, preferably dimeric, CD40-L can stimulate B cell proliferation. CD40 (receptor) antagonizes CD40-L proliferation of B cells.

Yet another assay for determining CD40-L biological activity is to measure immunoglobulin produced by B cells in response to activation by CD40-L or a derivative or analog thereof. Polyclonal immunoglobulin secretion can be measured, for example, by incubating with 5 x 10⁵ B cells/ml in culture for at least seven days. Immunoglobulin (Ig) production can be measured by an ELISA assay such as one described in Maliszewski et al., *J. Immunol.* 144:3028, 1990 [Maliszewski et al. I] or Maliszewski et al., *Eur J. Immunol.* 20:1735, 1990 [Maliszewski et al. II]. Murine B cells can be obtained, for example, from mice and cultured according to procedures described in Grabstein et al., *J. Exp. Med.* 163:1405, 1986 [Grabstein et al. I], Maliszewski et al. I, and Maliszewski et al. II.

CD40-L can be used in a binding assay to detect cells expressing CD40. For example, murine CD40-L according to Figure 1 (SEQ ID NO:1) or human CD40-L according to Figure 2 (SEQ ID NO:11), or an extracellular domain or a fragment thereof, can be conjugated to a detectable moiety such as ¹²⁵I. Radiolabeling with ¹²⁵I can be performed by any of several standard methodologies that yield a functional ¹²⁵I-CD40-L molecule labeled to high specific activity. Alternatively, another detectable moiety such as an enzyme that can catalyze a colorimetric or fluorometric reaction, biotin or avidin may be used. Cells expressing CD40 can be contacted with conjugated CD40-L. After incubation, unbound conjugated CD40-L is removed and binding is measured using the detectable moiety.

CD40-L polypeptides may exist as oligomers, such as dimers or trimers.

Oligomers are linked by disulfide bonds formed between cysteine residues on different CD40-L polypeptides. Alternatively, one can link two soluble CD40-L domains with a Gly4SerGly5Ser linker sequence, or other linker sequence described in United States

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Patent 5,073,627, which is incorporated by reference herein. CD40-L polypeptides may also be created by fusion of the C terminal of soluble CD40-L (extracellular domain) to the Fc region of IgG1 (for example, SEQ ID NO:3) as described for the CD40/Fc fusion protein. CD40-L/Fc fusion proteins are allowed to assemble much like heavy chains of an antibody molecule to form divalent CD40-L. If fusion proteins are made with both heavy and light chains of an antibody, it is possible to form a CD40-L oligomer with as many as four CD40-L extracellular regions.

Fusion proteins can be prepared using conventional techniques of enzyme cutting and ligation of fragments from desired sequences. PCR techniques employing synthetic oligonucleotides may be used to prepare and/or amplify the desired fragments. Overlapping synthetic oligonucleotides representing the desired sequences can also be used to prepare DNA constructs encoding fusion proteins. Fusion proteins can also comprise CD40-L and two or more additional sequences, including a leader (or signal peptide) sequence, Fc region, linker sequence, and sequences encoding highly antigenic moieties that provide a means for facile purification or rapid detection of a fusion protein.

Signal peptides facilitate secretion of proteins from cells. An exemplary signal peptide is the amino terminal 25 amino acids of the leader sequence of human interleukin-7 (IL-7; Goodwin et al., *Proc. Natl. Acad. Sci. U.S.A.* 86:302, 1989; Figure 2B). Other signal peptides may also be employed. For example, certain nucleotides in the IL-7 leader sequence can be altered without altering the amino acid sequence. Additionally, amino acid changes that do not affect the ability of the IL-7 sequence to act as a leader sequence can be made.

The Flag® octapeptide (Asp-Tyr-Lys-Asp-Asp-Asp-Asp-Lys) does not alter the biological activity of fusion proteins, is highly antigenic and provides an epitope reversibly bound by a specific monoclonal antibody, enabling rapid detection and facile purification of the expressed fusion protein. The Flag® sequence is also specifically cleaved by bovine mucosal enterokinase at the residue immediately following the Asp-Lys pairing, fusion proteins capped with this peptide may also be resistant to intracellular degradation in *E. coli*. A murine monoclonal antibody that binds the Flag® sequence has been deposited with the ATCC under accession number HB 9259; methods of using the antibody in purification of fusion proteins comprising the Flag® sequence are described in U.S. Patent 5,011,912, which is incorporated by reference herein.

Suitable Fc regions are defined as Fc regions that can bind to protein A or protein G, or alternatively, are recognized by an antibody that can be used in purification or detection of a fusion protein comprising the Fc region. Preferable Fc

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regions include the Fc region of human IgG₁ or murine IgG₁. One example is the human IgG₁ Fc region shown in SEQ ID NO:3; another example is an Fc region encoded by cDNA obtained by PCR from oligonucleotide primers from SEQ ID NO:9 and SEQ ID NO:10 with human cDNA as a template. Portions of a suitable Fc region may also be used, for example, an Fc region of human IgG₁ from which has been deleted a sequence of amino acids responsible for binding to protein A, such that the resultant Fc region binds to protein G but not protein A.

The [Gly4Ser]3 repeat sequence provides a linker sequence that separates the extracellular region of the CD40-L from the Fc portion of the fusion protein by a distance sufficient to ensure that the CD40-L properly folds into its secondary and tertiary structures. Suitable linker sequences (1) will adopt a flexible extended conformation, (2) will not exhibit a propensity for developing an ordered secondary structure which could interact with the functional domains of fusion proteins, and (3) will have minimal hydrophobic or charged character which could promote interaction with the functional protein domains. Typical surface amino acids in flexible protein regions include Gly, Asn and Ser. Virtually any permutation of amino acid sequences containing Gly, Asn and Ser would be expected to satisfy the above criteria for a linker sequence. Other near neutral amino acids, such as Thr and Ala, may also be used in the linker sequence. The length of the linker sequence may vary without significantly affecting the biological activity of the fusion protein. Linker sequences are unnecessary where the proteins being fused have non-essential N- or C-terminal amino acid regions which can be used to separate the functional domains and prevent steric interference.

CD40-L polypeptides may exist as soluble polypeptides comprising the extracellular domain of CD40-L as shown in Figure 1 (SEQ ID NO:1) and Figure 2 (SEQ ID NO:11) or as membrane-bound polypeptides comprising the extracellular domain, a transmembrane region and a short intracellular domain, as shown in Figure 1 (SEQ ID NO:1) and Figure 2 (SEQ ID NO:11) for the murine and human sequences, respectively. Moreover, the present invention comprises oligomers of CD40-L extracellular domains or fragments thereof, linked by disulfide interactions, or expressed as fusion polymers with or without spacer amino acid linking groups. For example, a dimer CD40-L molecule can be linked by an IgG Fc region linking group.

Without being bound by theory, membrane-bound CD40-L and oligomeric CD40-L can achieve activity stimulating Ig formation and proliferation of B cells previously only achieved by cross-linked anti-CD40 antibody in the presence of IL-4. It further appears likely that monomeric soluble CD40-L, comprising only the extracellular domain of CD40-L and capable of binding to CD40 receptor, will serve to antagonize the activity of membrane-bound and oligomeric CD40-L and/or cross-linked

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anti-CD40 antibodies. It further appears likely that the interaction of membrane-bound CD40-L with CD40 is the principal molecular interaction responsible for T cell contact dependent induction of B cell growth and differentiation to both antigen specific antibody production and polyclonal Ig secretion. In this regard, a mammalian cell transfected with a cDNA encoding full length CD40-L (i.e., being membrane-bound and having an intracellular domain, a transmembrane region and an extracellular domain or a fragment thereof) can mimic T cells in their ability to induce B cell growth, differentiation and stimulation of antigen-specific antibody production. It appears that activities of oligomeric soluble CD40-L, preferably a dimer of extracellular regions, can mimic the biological activities of membrane-bound CD40-L. Moreover, soluble monomeric CD40-L (comprising the extracellular domain or a fragment thereof) can bind to CD40 receptor to prevent T cell interaction with B cells and therefor have activity similar to CD40 (receptor) extracellular domain which itself may be in monomeric or in oligomeric form. Alternatively, CD40-L can be oligomeric (preferably a dimer) to act as a soluble factor capable of inducing B cell growth, differentiation and stimulation of antigen-specific antibody production. Accordingly, it appears that membrane-bound CD40-L and oligomeric CD40-L act as CD40 agonists, while soluble (monomeric) CD40-L and soluble CD40 act as CD40 antagonists by blocking CD40 receptor sites without significantly transducing signal or by preventing CD40-L binding to CD40 sites on B cells and other target cells.

Both CD40 agonists and CD40 antagonists will have useful therapeutic activity. For example, CD40 agonists (i.e., membrane-bound CD40-L and oligomeric CD40-L) are useful as vaccine adjuvants and for stimulating mAb production from hybridoma cells. CD40 antagonists (i.e., CD40 receptor, CD40/Fc and possibly soluble, monomeric CD40-L) are useful for treating autoimmune diseases characterized by presence of high levels of antigen-antibody complexes, such as allergy, lupus, rheumatoid arthritis, insulin dependent diabetes mellitus (IDDM), graft versus host disease (GVHD) and others.

IgE secretion from human B cells can be induced by IL-4 in the presence of T cells (Vercelli et al., *J. Exp. Med.* 169:1295, 1989). Further, IgE production can be induced from T cell depleted PBM (peripheral blood mononuclear cells) by addition of an anti-CD40 mAb (Jabara et al., *J. Exp. Med.* 172:1861, 1990 and Zhang et al., *J. Immunol.* 146:1836, 1991). The present invention further includes a method for inhibiting IgE production from activated B cells, activated by IL-4 in the presence of T cells or by CD40-L (preferably, membrane-bound CD40-L), comprising administering an effective amount of a CD40/Fc fusion protein, as described herein, or a soluble CD40 encoded by the cDNA sequence described in SEQ ID NO. 3. Similarly, CD40

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receptors and possibly soluble CD40-L (monomer only) can also block secretion of other antibody isotypes.

The present invention further includes CD40-L polypeptides with or without associated native-pattern glycosylation. CD40-L expressed in yeast or mammalian expression systems (e.g., COS-7 cells) may be similar to or significantly different from a native CD40-L polypeptide in molecular weight and glycosylation pattern, depending upon the choice of expression system. Expression of CD40-L polypeptides in bacterial expression systems, such as *E. coli*, provides non-glycosylated molecules.

DNA constructs that encode various additions or substitutions of amino acid residues or sequences, or deletions of terminal or internal residues or sequences not needed for biological activity or binding can be prepared. For example, the extracellular CD40-L N-glycosylation site can be modified to preclude glycosylation while allowing expression of a homogeneous, reduced carbohydrate analog using yeast expression systems. N-glycosylation sites in eukaryotic polypeptides are characterized by an amino acid triplet Asn-X-Y, wherein X is any amino acid except Pro and Y is Ser or Thr. Appropriate modifications to the nucleotide sequence encoding this triplet will result in substitutions, additions or deletions that prevent attachment of carbohydrate residues at the Asn side chain. In another example, sequences encoding Cys residues can be altered to cause the Cys residues to be deleted or replaced with other amino acids, preventing formation of incorrect intramolecular disulfide bridges upon renaturation. Human CD40-L comprises five Cys residues in its extracellular domain. Thus, at least one of the five Cys residues can be replaced with another amino acid or deleted without effecting protein tertiary structure or disulfide bond formation.

Other approaches to mutagenesis involve modification of sequences encoding dibasic amino acid residues to enhance expression in yeast systems in which KEX2 protease activity is present. Sub-units of a CD40-L polypeptide may be constructed by deleting sequences encoding terminal or internal residues or sequences.

CD40-L polypeptides are encoded by multi-exon genes. The present invention further includes alternative mRNA constructs which can be attributed to different mRNA splicing events following transcription and which share regions of identity or similarity with the cDNAs disclosed herein.

Antisense or sense oligonucleotides comprise a single-stranded nucleic acid sequence (either RNA or DNA) capable of binding to target CD40-L mRNA (sense) or CD40-L DNA (antisense) sequences. Antisense or sense oligonucleotides, according to the present invention, comprise a fragment of SEQ ID NO:1 or SEQ ID NO:11, or a DNA or RNA complement of SEQ ID NO:1 or SEQ ID NO:11. Such a fragment comprises at least about 14 nucleotides. Preferably, such a fragment comprises from

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about 14 to about 30 nucleotides. The ability to create an antisense or a sense oligonucleotide, based upon a cDNA sequence for CD40-L is described in, for example, Stein and Cohen, *Cancer Res.* 48:2659, 1988 and van der Krol et al., *BioTechniques* 6:958, 1988.

Binding of antisense or sense oligonucleotides to target nucleic acid sequences results in the formation of duplexes that block translation (RNA) or transcription (DNA) by one of several means, including enhanced degradation of the duplexes, premature termination of transcription or translation, or by other means. Suitable polymerase promotors include promotors for any RNA polymerase, or promotors for any DNA polymerase. Antisense or sense oligonucleotides further comprise oligonucleotides having modified sugar-phosphodiester backbones (or other sugar linkages, such as those described in WO91/06629) and wherein such sugar linkages are resistant to endogenous nucleases. Such oligonucleotides with resistant sugar linkages are stable in vivo (i.e., capable of resisting enzymatic degradation) but retain sequence specificity to be able to bind to target nucleotide sequences. Other examples of sense or antisense oligonucleotides include those oligonucleotides which are covalently linked to organic moieties, such as those described in WO 90/10448, and other moieties that increases affinity of the oligonucleotide for a target nucleic acid sequence, such as poly-(L-lysine). Further still, intercalating agents, such as ellipticine, and alkylating agents or metal complexes may be attached to sense or antisense oligonucleotides to modify binding specificities of the antisense or sense oligonucleotide for the target nucleotide sequence. Antisense or sense oligonucleotides may be introduced into a cell containing the target nucleic acid sequence by any gene transfer method, including, for example, CaPO₄-mediated DNA transfection, electroporation, or other gene transfer vectors such as Epstein-Barr virus. Antisense or sense oligonucleotides are preferably introduced into a cell containing the target nucleic acid sequence by insertion of the antisense or sense oligonucleotide into a suitable retroviral vector, then contacting the cell with the retrovirus vector containing the inserted sequence, either in vivo or ex vivo. Suitable retroviral vectors include, but are not limited to, the murine retrovirus M-MuLV, N2 (a retrovirus derived from M-MuLV), or or the double copy vectors designated DCT5A, DCT5B and DCT5C (see PCT Application US 90/02656). Alternatively, other promotor sequences may be used to express the oligonucleotide.

Sense or antisense oligonucleotides may also be introduced into a cell containing the target nucleotide sequence by formation of a conjugate with a ligand binding molecule, as described in WO 91/04753. Suitable ligand binding molecules include, but are not limited to, cell surface receptors, growth factors, other cytokines, or other ligands that bind to cell surface receptors. Preferably, conjugation of the

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ligand binding molecule does not substantially interfere with the ability of the ligand binding molecule to bind to its corresponding molecule or receptor, or block entry of the sense or antisense oligonucleotide or its conjugated version into the cell.

Alternatively, a sense or an antisense oligonucleotide may be introduced into a cell containing the target nucleic acid sequence by formation of an oligonucleotide-lipid complex, as described in WO 90/10448. The sense or antisense oligonucleotide-lipid complex is preferably dissociated within the cell by an endogenous lipase.

The sequence of murine CD40-L cDNA was obtained by direct expression techniques. The sequence of human CD40-L was obtained by cross-species hybridization techniques using the murine CD40-L cDNA as a probe.

We cloned murine CD40-L by first obtaining a clone of the extracellular region of human CD40 (the receptor) by polymerase chain reaction (PCR) techniques using primers based upon a sequence published in Stamenkovic et al. (SEQ ID NO:4). An upstream oligonucleotide primer 5'-CCGTCGACCACCATGGTTCGTCTGCC -3' (SEQ ID NO:5) introduces a Sal 1 site upstream from an initiator methionine of CD40 and a downstream oligonucleotide primer 5'-CCGTCGACGTCTAGAGCCGATCCTGGGG-3' (SEQ ID NO:6) inserts a termination codon after amino acid 192 of CD40, followed by Xba 1 and Sal 1 sites. The amplified cDNA was digested with Sal 1 and cloned into pDC406 (McMahan et al., EMBO J. 10:2821, 1991) to construct pDC406/s CD40.

A second CD40 receptor fragment (SEQ ID NO:4) was obtained by PCR techniques for fusion to the Fc domain of human IgG1 (SEQ ID NO:3). Briefly, The upstream oligonucleotide primer (SEQ ID NO:5) and fusion template (SEQ ID NO:4) were the same as before. The downstream oligonucleotide primer was 5'-

ACAAGATCTGGGCTCTACGTATCTCAGCCGATCCTGGGGAC-3' (SEQ ID NO:7) that inserts amino acids Tyr Val Glu Pro Arg (SEQ ID NO:8) after amino acid 193 of CD40. Glu and Pro are the first two amino acids of a hinge region of human IgG1, and are followed by a Bgl II restriction site. The Bgl II restriction site was used to fuse the extracellular domain of CD40 to the remainder of human IgG1 Fc region.

Other fusion proteins comprising ligand binding domains from other receptors can be made by obtaining a DNA sequence for the ligand binding domain of a receptor and fusing this sequence to a DNA sequence encoding an Fc region of an antibody molecule that binds to protein A or protein G, or another polypeptide that is capable of affinity purification, for example, avidin or streptavidin. The resultant gene construct can be introduced into mammalian cells to transiently express a fusion protein. Receptor/Fc fusion proteins can be purified by protein A or protein G affinity purification. Receptor/avidin fusion proteins can be purified by biotin affinity

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chromatography. The fusion protein can later be removed from the column by eluting with a high salt solution or another appropriate buffer.

We obtained a cDNA encoding human IgG1 Fc region by PCR amplification using cDNA from human cells as a template and an upstream oligonucleotide primer 5'-TATTAATCATTCAGTAGGGCCCAGATCTTGTGACAAAACTCAC-3' (SEQ ID 5 NO:9) and a downstream oligonucleotide primer 5'-GCCAGCTTAACTAGTTCATTTACCCGGAGACAGGGAGA-3" (SEQ ID NO:10). The PCR amplified cDNA introduced a Bgl II site near the beginning of the hinge region, which was used to ligate CD40 extracellular domain to construct a s CD40/Fc fusion cDNA, which was ligated into pDC406 to construct pDC406/CD40/Fc. Other 10 suitable Fc regions are defined as any region that can bind with high affinity to protein A or protein G, and includes the Fc region of human IgG1 or murine IgG1. One example is the human IgG1 Fc region shown in SEQ ID NO:3 or the cDNA obtained by PCR from oligonucleotide primers from SEQ ID NO:9 and SEQ ID NO:10 with 15 human cDNA as a template.

Receptor/Fc fusion molecules preferably are synthesized in recombinant mammalian cell culture because they are generally too large and complex to be synthesized by prokaryotic expression methods. Examples of suitable mammalian cells for expressing a receptor/Fc fusion protein include CV-1 cells (ATCC CCL 70) and COS-7 cells (ATCC CRL 1651), both derived from monkey kidney.

The DNA construct pDC406/CD40/Fc was transfected into the monkey kidney cell line CV-1/EBNA (ATCC CRL 10478). The pDC406 plasmid includes regulatory sequences derived from SV40, human immunodeficiency virus (HIV), and Epstein-Barr virus (EBV). The CV-1/EBNA cell line was derived by transfection of the CV-1 cell line with a gene encoding Epstein-Barr virus nuclear antigen-1 (EBNA-1) and constitutively express EBNA-1 driven from human CMV immediate-early enhancer/promoter. An EBNA-1 gene allows for episomal replication of expression vectors, such as pDC406, that contain the EBV origin of replication.

Transfectants expressing CD40/Fc fusion protein are initially identified using dot blots or Western blots. The supernatants are then subjected to dot blot or gel electrophoresis followed by transfer of the electrophoresed proteins for binding to G28-5 mAb (an antibody that binds to human CD40 receptor). The blotted proteins were then incubated with radiolabeled with ¹²⁵I-protein A, washed to remove unbound label, and examined for expression of Fc. Monoclonal antibody G28-5 was produced according to Clark et al., *supra*.

Once cells expressing the fusion construct were identified, large scale cultures of transfected cells were grown to accumulate supernatant from cells expressing

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CD40/Fc. CD40/Fc fusion protein in supernatant fluid was purified by affinity purification. Briefly, one liter of culture supernatant containing CD40/Fc fusion protein was purified by filtering mammalian cell supernatants (e.g., in a 0.45µ filter) and applying filtrate to a protein A/G antibody affinity column (Schleicher and Schuell, Keene, NH) at 4°C at a flow rate of 80 ml/hr for a 1.5 cm x 12.0 cm column. The

column was washed with 0.5 M NaCl in PBS until free protein could not be detected in wash buffer. Finally, the column was washed with PBS. Bound fusion protein was eluted from the column with 25 mM citrate buffer, pH 2.8, and brought to pH 7 with 500 mM Hepes buffer, pH 9.1. Silver-stained SDS gels of the eluted CD40/Fc fusion protein showed it to be > 98% pure.

Soluble CD40 (sCD40) and CD40/Fc fusion proteins were made as described herein. The supernatants were purified through a G28-5 (anti-CD40 mAb) affinity column to affinity purify sCD40 expressed by the transfected CV-1/EBNA cells. Protein-containing fractions were pooled and aliquots removed for G28-5 binding assays and analysis by SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis) in the presence of 1 mM dithiothreitol as a reducing agent. A single band was seen of molecular weight 28,100 daltons. In the absence of a reducing agent, SDS-PAGE analysis of sCD40 revealed two bands, a major band of molecular weight 56,000 and a minor band of molecular weight 28,000. The banding pattern indicates that the majority of sCD40 exists as a disulfide-linked homodimer in solution. The 28,000 band is free monomer.

CD40 proteins were visualized by silver staining. Sample protein concentrations were determined using a micro-BCA assay (Pierce) with ultrapure bovine serum albumin as standard. Soluble CD40 purity and protein concentration were confirmed by amino acid analysis. Purified soluble CD40 was absorbed to PVDF paper and the paper subjected to automated Edman degradation on an Applied Biosystems model 477A protein sequencer according to manufacturers instructions for N-terminal protein sequencing. This procedure checked the protein sequence of sCD40.

30 Soluble CD40 and CD40/Fc fusion protein were able to modulate human B cell responses in the absence of anti-CD40 mAb (G28-5). Purified tonsillar B cells were cultured with anti-IgM and human IL-4 and either sCD40 or CD40/Fc fusion protein was added. Neither form of CD40 had an inhibitory effect on B cell proliferation (as measured by tritiated thymidine incorporation). IL-4 receptor, by contrast, inhibited IL-4-induced B cell proliferation in a concentration-dependent manner.

Soluble CD40 and CD40/Fc were tested for their ability to inhibit IL-4 induced IgE secretion in a 2-donor MLC (mixed lymphocyte culture) system. In three

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experiments, the level of IgE production was reduced as the concentration of CD40 was increased. Soluble CD40, added at a concentration of 10 µg/ml, was able to completely inhibit IgE secretion in this model of allergy. Further, CD40/Fc had similar effects as its soluble counterpart. However, addition of an IL-7 receptor-Fc fusion protein (made by similar procedures with a published IL-7 receptor sequence) did not affect secretion of IgE in this model.

Levels of CD23 were also measured in the same MLC in response to sCD40 or CD40/Fc fusion proteins. Soluble CD40 produced a small, but reproducible decrease in sCD23 level at day 6 compared to cultures stimulated with IL-4 alone, however a stronger inhibitory effect was pronounced at day 12 in the same cultures. Soluble CD23 induction by IL-4-stimulated T-depleted PBM (peripheral blood macrophages) E-cells was similarly affected by addition of sCD40, causing a small decrease in sCD23 levels at day 6 and a more pronounced inhibition at day 12. In each culture system, the results with CD40/Fc fusion protein were substantially the same as with sCD40.

In an effort to isolate a cDNA for a CD40-L, purified CD40/Fc fusion protein was radioiodinated with ¹²⁵I using a commercially available solid phase agent (IODO-GEN, Pierce). In this procedure, 5 µg of IODO-GEN were plated at the bottom of a 10 x 75 mm glass tube and incubated for twenty minutes at 4° C with 75 µl of 0.1 M sodium phosphate, pH 7.4 and 20 µl (2 mCi) Na¹²⁵I. The solution was then transferred to a second glass tube containing 5 µg of CD40/Fc in 45 µl PBS (phosphate buffered saline) and this reaction mixture was incubated for twenty minutes at 4° C. The reaction mixture was fractionated by gel filtration on a 2 ml bed volume of Sephadex® G-25 (Sigma), and then equilibrated in RPMI 1640 medium containing 2.5% (v/v) bovine serum albumin (BSA), 0.2% (v/v) sodium azide and 20 mM Hepes, pH 7.4 binding medium. The final pool of ¹²⁵I CD40/Fc was diluted to a working stock solution of 1 x 10⁻⁷ M in binding medium and stored for up to one month at 4° C without detectable loss of receptor binding activity.

A cDNA library was prepared from a EL4 cell line sorted by FACS (fluorescence activated cell sorting) on the basis of binding of a biotinylated CD40/Fc fusion protein. Cells were sorted five times until there was a significant shift in fluorescence intensity based upon expression of a ligand for CD40 by the sorted EL-4 cells. The five-times sorted cells were called EL-40.5 cells and these cells were cultured for the purposes of creating a cDNA library from EL-40.5 mRNA. Briefly, cDNA was synthesized, inserted into empty pDC406 vector and transformed into *E. coli*. Transformants were pooled, and the DNA from the pools was isolated and transfected into CV1-EBNA cells to create an expression cloning library. Transfected CV1-EBNA cells were cultured on slides for three days to permit transient expression

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of CD40-L. The slides containing the transfected cells were then incubated with radioiodinated CD40/Fc, washed to remove unbound CD40/Fc, and fixed with gluteraldehyde. The fixed slides were dipped in liquid photographic emulsion and exposed in the dark. After developing the slides, they were individually examined with a microscope and cells expressing CD40-L were identified by the presence of autoradiographic silver grains against a light background.

The expression cloning library from EL-40.5 cells was screened and one pool, containing approximately 2000 individual clones, was identified as positive for binding ¹²⁵I labeled CD40/Fc fusion protein. This pool was broken down into smaller pools of approximately 200 colonies. The smaller pools were screened as described above. One of the smaller pools was positive for CD40-L.

A single clone was isolated and sequenced by standard techniques, to provide the cDNA sequence and deduced amino acid sequence of murine CD40-L as shown in Figure 1 and SEQ ID NO:1.

15 The human homolog CD40-L cDNA was found by cross species hybridization techniques. Briefly, a human peripheral blood lymphocyte (PBL) cDNA library was made from peripheral blood lymphocytes treated with OKT3 antibody (ATCC, Rockville MD) that binds to CD3 (10 ng/ml) and interleukin-2 (IL-2, 10 ng/ml) for six days. The PBL cells were washed and then stimulated for 4 hours with 10 ng/ml PMA 20 (phorbol myristate acetate, Sigma St Louis) and 500 ng/ml ionomycin (Calbiochem). Messenger RNA was isolated from stimulated PBL cells, cDNA formed and cDNA was ligated into Eco R1 linkers. Ligated cDNA was inserted into the Eco R1 site of λgt10 phage cloning vehicle (Gigapak® Stratagene, San Diego, CA) according to manufacturer's instructions. Phage were amplified, plated at densities densities of 25 approximately 20,000 phage per 15 cm plate. and phage lifts were performed, as described in Maniatis et al., Molecular Biology: A Laboratory Manual, Cold Spring Harbor Laboratory, NY, 1982, pages 316-328. A murine probe was constructed corresponding to the coding region of murine CD40-L from nucleotide 13 to nucleotide 793 of SEQ ID NO:1 and Figure 1. This probe was hybridized to to the PBL library 30 phage lifts under conditions of moderate to severe stringency. Briefly, hybridization conditions were 6 X SSC, 1 X Denhardt's solution, 2 mM EDTA, 0.5% Np40 (Nonidet P-40 detergent) at 63 °C overnight. This was followed by washing in 3 X SSC, 0.1% SDS for three hours at 55°C, followed by overnight exposure to X-Ray film. Positive plaques were identified at a frequency of approximately 1 per 1000 35 plaques. Positive plaques were purified twice and cDNA was prepared from amplified cultures.

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One can utilize the murine or human CD40-L cDNA sequences disclosed herein to obtain cDNAs encoding other mammalian homologs of murine or human CD40-L by cross-species hybridization techniques. Briefly, an oligonucleotide probe is created from the nucleotide sequence of the extracellular region of murine CD40-L as described in Figure 1 (SEQ ID NO:1) or human CD40-L as described in Figure 2 (SEQ ID NO:11). This probe can be made by standard techniques, such as those described in Maniatis et al. *supra*. The murine or human probe is used to screen a mammalian cDNA library or genomic library under moderate stringency conditions. Examples of mammalian cDNA or genomic libraries include, for cDNA, a library made from the mammal's peripheral blood lymphocytes. Alternatively, various cDNA libraries or mRNAs isolated from various cell lines can be screened by Northern hybridization to determine a suitable source of mammalian CD40-L DNA or mRNA.

Recombinant expression vectors for expression of CD40-L by recombinant DNA techniques include a CD40-L DNA sequence comprising a synthetic or cDNAderived DNA fragment encoding a CD40-L polypeptide, operably linked to a suitable 15 transcriptional or translational regulatory nucleotide sequence, such as one derived from a mammalian, microbial, viral, or insect gene. Examples of regulatory sequences include sequences having a regulatory role in gene expression (e.g., a transcriptional promoter or enhancer), optionally an operator sequence to control transcription, a sequence encoding an mRNA ribosomal binding site, and appropriate sequences which 20 control transcription and translation initiation and termination. Nucleotide sequences are operably linked when the regulatory sequence functionally relates to the CD40-L DNA sequence. Thus, a promoter nucleotide sequence is operably linked to a CD40-L DNA sequence if the promoter nucleotide sequence controls the transcription of the CD40-L DNA sequence. Still further, a ribosome binding site may be operably linked 25 to a sequence for a CD40-L polypeptide if the ribosome binding site is positioned within the vector to encourage translation. In addition, sequences encoding signal peptides can be incorporated into expression vectors. For example, a DNA sequence for a signal peptide (secretory leader) may be operably linked to a CD40-L DNA sequence. The signal peptide is expressed as a precursor amino acid sequence which 30 enables improved extracellular secretion of translated fusion polypeptide by a yeast host cell.

Suitable host cells for expression of CD40-L polypeptides include prokaryotes, yeast or higher eukaryotic cells. Prokaryotes include gram negative or gram positive organisms, for example, *E. coli* or *Bacilli*. Suitable prokaryotic host cells for transformation include, for example, *E. coli*, *Bacillus subtilis*, *Salmonella typhimurium*, and various other species within the genera *Pseudomonas*,

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Streptomyces, and Staphylococcus. Higher eukaryotic cells include established cell lines of mammalian origin. Cell-free translation systems could also be employed to produce CD40-L polypeptides using RNAs derived from DNA constructs disclosed herein. Appropriate cloning and expression vectors for use with bacterial, fungal, yeast, and mammalian cellular hosts are described, for example, in Pouwels et al. Cloning Vectors: A Laboratory Manual, Elsevier, New York, (1985).

In a prokaryotic host cell, such as *E. coli*, a CD40-L polypeptide or analog may include an N-terminal methionine residue to facilitate expression of the recombinant polypeptide in the prokaryotic host cell. The N-terminal Met may be cleaved from the expressed recombinant CD40-L polypeptide. Prokaryotic host cells may be used for expression of CD40-L polypeptides that do not require extensive proteolytic or disulfide processing.

The expression vectors carrying the recombinant CD40-L DNA sequence are transfected or transformed into a substantially homogeneous culture of a suitable host microorganism or mammalian cell line. Transformed host cells are cells which have been transformed or transfected with nucleotide sequences encoding CD40-L polypeptides and express CD40-L polypeptides. Expressed CD40-L polypeptides will be located within the host cell and/or secreted into culture supernatant fluid, depending upon the nature of the host cell and the gene construct inserted into the host cell.

Expression vectors transfected into prokaryotic host cells generally comprise one or more phenotypic selectable markers. A phenotypic selectable marker is, for example, a gene encoding a protein that confers antibiotic resistance or that supplies an autotrophic requirement, and an origin of replication recognized by the host to ensure amplification within the host. Other useful expression vectors for prokaryotic host cells include a selectable marker of bacterial origin derived from commercially available plasmids. This selectable marker can comprise genetic elements of the cloning vector pBR322 (ATCC 37017). pBR322 contains genes for ampicillin and tetracycline resistance and thus provides simple means for identifying transformed cells. The pBR322 "backbone" sections are combined with an appropriate promoter and a CD40-L DNA sequence. Other commercially vectors include, for example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and pGEM1 (Promega Biotec, Madison, WI, USA).

Promoter sequences are commonly used for recombinant prokaryotic host cell expression vectors. Common promoter sequences include β-lactamase (penicillinase), lactose promoter system (Chang et al., *Nature 275:615*, 1978; and Goeddel et al., *Nature 281:544*, 1979), tryptophan (trp) promoter system (Goeddel et al., *Nucl. Acids Res. 8:4057*, 1980; and EP-A-36776) and tac promoter (Maniatis, *Molecular Cloning:*

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A Laboratory Manual, Cold Spring Harbor Laboratory, p. 412, 1982). A particularly useful prokaryotic host cell expression system employs a phage λ P_L promoter and a cI857ts thermolabile repressor sequence. Plasmid vectors available from the American Type Culture Collection which incorporate derivatives of the λ P_L promoter include plasmid pHUB2 (resident in *E. coli* strain JMB9 (ATCC 37092)) and pPLc28 (resident in *E. coli* RR1 (ATCC 53082)).

CD40-L may be expressed in yeast host cells, preferably from the Saccharomyces genus (e.g., S. cerevisiae). Other genera of yeast, such as Pichia or Kluyveromyces, may also be employed. Yeast vectors will often contain an origin of replication sequence from a 2µ yeast plasmid, an autonomously replicating sequence (ARS), a promoter region, sequences for polyadenylation, and sequences for transcription termination. Preferably, yeast vectors include an origin of replication sequence and selectable marker. Suitable promoter sequences for yeast vectors include promoters for metallothionein, 3-phosphoglycerate kinase (Hitzeman et al., J. Biol. Chem. 255:2073, 1980) or other glycolytic enzymes (Hess et al., J. Adv. Enzyme Reg. 7:149, 1968; and Holland et al., Biochem. 17:4900, 1978), such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase. Other suitable vectors and promoters for use in yeast expression are further described in Hitzeman, EPA-73,657.

Yeast vectors can be assembled, for example, using DNA sequences from pBR322 for selection and replication in *E. coli* (Amp^r gene and origin of replication). Other yeast DNA sequences that can be included in a yeast expression construct include a glucose-repressible ADH2 promoter and α-factor secretion leader. The ADH2 promoter has been described by Russell et al. (*J. Biol. Chem. 258:2674*, 1982) and Beier et al. (*Nature 300:*724, 1982). The yeast α-factor leader sequence directs secretion of heterologous polypeptides. The α-factor leader sequence is often inserted between the promoter sequence and the structural gene sequence. *See, e.g.*, Kurjan et al., *Cell 30:*933, 1982 and Bitter et al., *Proc. Natl. Acad. Sci. USA 81:*5330, 1984. Other leader sequences suitable for facilitating secretion of recombinant polypeptides from yeast hosts are known to those of skill in the art. A leader sequence may be modified near its 3' end to contain one or more restriction sites. This will facilitate fusion of the leader sequence to the structural gene.

Yeast transformation protocols are known to those of skill in the art. One such protocol is described by Hinnen et al., *Proc. Natl. Acad. Sci. USA 75*:1929, 1978. The Hinnen et al. protocol selects for Trp⁺ transformants in a selective medium,

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wherein the selective medium consists of 0.67% yeast nitrogen base, 0.5% casamino acids, 2% glucose, 10 μ g/ml adenine and 20 μ g/ml uracil.

Yeast host cells transformed by vectors containing ADH2 promoter sequence may be grown for inducing expression in a "rich" medium. An example of a rich medium is one consisting of 1% yeast extract, 2% peptone, and 1% glucose supplemented with 80 μ g/ml adenine and 80 μ g/ml uracil. Derepression of the ADH2 promoter occurs when glucose is exhausted from the medium.

Mammalian or insect host cell culture systems could also be employed to express recombinant CD40-L polypeptides. Examples of suitable mammalian host cell lines include the COS-7 line of monkey kidney cells (ATCC CRL 1651) (Gluzman et al., Cell 23:175, 1981), L cells, C127 cells, 3T3 cells (ATCC CCL 163), Chinese hamster ovary (CHO) cells, HeLa cells, and BHK (ATCC CRL 10) cell lines. Suitable mammalian expression vectors include nontranscribed elements such as an origin of replication, a promoter sequence, an enhancer linked to the structural gene, other 5' or 3' flanking nontranscribed sequences, such as ribosome binding sites, a polyadenylation site, splice donor and acceptor sites, and transcriptional termination sequences.

Transcriptional and translational control sequences for mammalian host cell expression vectors may be excised from viral genomes. For example, commonly used 20 mammalian cell promoter sequences and enhancer sequences are derived from Polyoma virus, Adenovirus 2, Simian Virus 40 (SV40), and human cytomegalovirus. DNA sequences derived from the SV40 viral genome, for example, SV40 origin, early and late promoter, enhancer, splice, and polyadenylation sites may be used to provide the other genetic elements required for expression of a structural gene sequence in a 25 mammalian host cell. Viral early and late promoters are particularly useful because both are easily obtained from a viral genome as a fragment which may also contain a viral origin of replication (Fiers et al., Nature 273:113, 1978). Smaller or larger SV40 fragments may also be used, provided the approximately 250 bp sequence extending from the Hind III site toward the Bgl I site located in the SV40 viral origin of 30 replication site is included.

Exemplary mammalian expression vectors can be constructed as disclosed by Okayama and Berg (*Mol. Cell. Biol. 3*:280, 1983). A useful high expression vector, PMLSV N1/N4, described by Cosman et al., *Nature 312*:768, 1984 has been deposited as ATCC 39890. Additional useful mammalian expression vectors are described in EP-A-0367566, and in U.S. Patent Application Serial No. 07/701,415, filed May 16, 1991, incorporated by reference herein. For expression of a type II protein extracellular region, such as CD40-L, a heterologous signal sequence should be added,

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such as the signal sequence for interleukin-7 (IL-7) described in United States Patent 4,965,195, or the signal sequence for interleukin-2 receptor described in United States Patent Application 06/626,667 filed on July 2, 1984.

Human or murine CD40-L can be made in membrane-bound form when an intracellular and transmembrane regions are included or in soluble form with only the extracellular domain. We expressed full length murine CD40-L in mammalian cells to yield cells expressing membrane-bound murine CD40-L. CV1 cells were transfected with a cDNA shown in Figure 1 (SEO ID NO:1) in HAVEO vector or CV1 cells were transfected with HAVEO empty vector using techniques described in Example 6 herein. This yielded transfected CV1 cells expressing membrane-bound murine CD40-L. These cells were used as a source of membrane-bound murine CD40-L for the series of experiments reported in Examples 10 -13 reported below.

Purification of Recombinant CD40-L Polypeptides

CD40-L polypeptides may be prepared by culturing transformed host cells under culture conditions necessary to express CD40-L polypeptides. The resulting expressed polypeptides may then be purified from culture media or cell extracts. A CD40-L polypeptide, if desired, may be concentrated using a commercially available protein concentration filter, for example, an Amicon or Millipore Pellicon ultrafiltration unit. Following the concentration step, the concentrate can be applied to a purification matrix such as a gel filtration medium. Alternatively, an anion exchange resin can be employed, for example, a matrix or substrate having pendant diethylaminoethyl (DEAE) groups. The matrices can be acrylamide, agarose, dextran, cellulose or other types commonly employed in protein purification. Alternatively, a cation exchange step can be employed. Suitable cation exchangers include various insoluble matrices comprising sulfopropyl or carboxymethyl groups. Sulfopropyl groups are preferred.

Finally, one or more reverse-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media, (e.g., silica gel having pendant methyl or other aliphatic groups) can be employed to further purify CD40-L. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a substantially homogeneous recombinant protein.

It is also possible to utilize an affinity column comprising CD40 ligand binding domain to affinity-purify expressed CD40-L polypeptides. CD40-L polypeptides can be removed from an affinity column in a high salt elution buffer and then dialyzed into a lower salt buffer for use.

Recombinant protein produced in bacterial culture is usually isolated by initial disruption of the host cells, centrifugation, extraction from cell pellets if an insoluble polypeptide, or from the supernatant fluid if a soluble polypeptide, followed by one or

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more concentration, salting-out, ion exchange, affinity purification or size exclusion chromatography steps. Finally, RP-HPLC can be employed for final purification steps. Microbial cells can be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents.

Transformed yeast host cells are preferably employed to express CD40-L as a secreted polypeptide. This simplifies purification. Secreted recombinant polypeptide from a yeast host cell fermentation can be purified by methods analogous to those disclosed by Urdal et al. (*J. Chromatog.* 296:171, 1984). Urdal et al. describe two sequential, reversed-phase HPLC steps for purification of recombinant human IL-2 on a preparative HPLC column.

Administration of CD40-L Compositions

The present invention provides therapeutic compositions comprising an effective amount of CD40-L in a suitable diluent or carrier and methods of treating mammals using the compositions. For therapeutic use, purified CD40-L or a biologically active analog thereof is administered to a patient, preferably a human, for treatment in a manner appropriate to the indication. Thus, for example, CD40-L pharmaceutical compositions (for example, in the form of a soluble extracellular domain, or a fragment thereof) which is administered to achieve a desired therapeutic effect can be given by bolus injection, continuous infusion, sustained release from implants, or other suitable technique. Typically, a CD40-L therapeutic agent will be administered in the form of a pharmaceutical composition comprising purified CD40-L polypeptide in conjunction with physiologically acceptable carriers, excipients or diluents. Such carriers will be nontoxic to patients at the dosages and concentrations employed. Ordinarily, the preparation of such compositions entails combining a CD40-L polypeptide with buffers, antioxidants such as ascorbic acid, low molecular weight (less than about 10 residues) polypeptides, proteins, amino acids, carbohydrates including glucose, sucrose or dextrans, chelating agents such as EDTA, glutathione and other stabilizers and excipients. Neutral buffered saline or saline mixed with conspecific serum albumin are exemplary appropriate diluents. CD40-L sense or antisense oligonucleotides may be administered in vivo by administering an effective amount of a vector containing a nucleic acid sequence that encodes and effective antisense or sense oligonucleotide. Additionally, CD40-L sense or antisense oligonucleotides may be administered ex vivo by removing cells containing CD40-L DNA or mRNA from an individual, incorporating an antisense or sense oligonucleotide into the cells using gene transfer techniques, and re-infusing the cells into the individual.

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The following examples are intended to illustrate particular embodiments and not limit the scope of the invention.

EXAMPLE 1

This example describes construction of a CD40/Fc DNA construct to express a soluble CD40/Fc fusion protein for use in detecting cDNA clones encoding a CD40 ligand. The cDNA sequence of the extracellular region or ligand binding domain of complete CD40 human receptor sequence was obtained using polymerase chain reaction (PCR) techniques, and is based upon the sequence published in Stamenkovic et al., supra. A CD40 plasmid (CDM8) was used as a template for PCR amplification. CDM8 is described in Stamenkovic et al. and was obtained from the authors. A PCR technique (Sarki et al., Science 239:487, 1988) was employed using 5' (upstream) and 3' (downstream) oligonucleotide primers to amplify the DNA sequences encoding CD40 extracellular ligand binding domain. Upstream oligonucleotide primer 5'-CCGTCGACCACCATGGTTCGTCTGCC -3' (SEQ ID NO:5) introduces a Sal 1 site upstream from an initiator methionine of CD40 and a downstream oligonucleotide primer 5'-ACAAGATCTGGGCTCTACGTATCTCAGCCGATCCTGGGGAC-3' (SEQ ID NO:7) that inserts amino acids Tyr Val Glu Pro Arg (SEQ ID NO:8) after amino acid 193 of CD40. Glu and Pro are the first two amino acids of a hinge region of human IgG1, and are followed by a $Bgl \, \Pi$ restriction site that was used to fuse the extracellular domain of CD40 to the remained of human IgG1 Fc region.

The DNA construct pDC406/CD40/Fc was transfected into the monkey kidney cell line CV-1/EBNA (ATCC CRL 10478). The pDC406 plasmid includes regulatory sequences derived from SV40, human immunodeficiency virus (HIV), and Epstein-Barr virus (EBV). The CV-1/EBNA cell line was derived by transfection of the CV-1 cell line with a gene encoding Epstein-Barr virus nuclear antigen-1 (EBNA-1) that constitutively expresses EBNA-1 driven from the human CMV intermediate-early enhancer/promoter. The EBNA-1 gene allows for episomal replication of expression vectors, such as pDC406, that contain the EBV origin of replication.

Once cells expressing the fusion construct were identified, large scale cultures of transfected cells were grown to accumulate supernatant from cells expressing CD40/Fc. The CD40/Fc fusion protein in supernatant fluid was purified by affinity purification. Briefly, one liter of culture supernatant containing the CD40/Fc fusion protein was purified by filtering mammalian cell supernatants (e.g., in a 0.45µ filter) and applying filtrate to a protein A/G antibody affinity column (Schleicher and Schuell, Keene, NH) at 4°C at a flow rate of 80 ml/hr for a 1.5 cm x 12.0 cm column. The column was washed with 0.5 M NaCl in PBS (phosphate buffered saline) until free

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protein could not be detected in wash buffer. Finally, the column was washed with PBS. Bound fusion protein was eluted from the column with 25 mM citrate buffer, pH 2.8, and brought to pH 7 with 500 mM Hepes buffer, pH 9.1. Silver-stained SDS gels of the eluted CD40/Fc fusion protein showed it to be > 98% pure.

Purified CD40/Fc fusion protein was iodinated with ¹²⁵I using a commercially available solid phase agent (IODO-GEN, Pierce). In this procedure, 5 μg of IODO-GEN were plated at the bottom of a 10 x 75 mm glass tube and incubated for twenty minutes at 4° C with 75 μl of 0.1 M sodium phosphate, pH 7.4 and 20 μl (2 mCi) Na¹²⁵I. The solution was then transferred to a second glass tube containing 5 μg of CD40/Fc in 45 μl PBS and this reaction mixture was incubated for twenty minutes at 4° C. The reaction mixture was fractionated by gel filtration on a 2 ml bed volume of Sephadex[®] G-25 (Sigma), and then equilibrated in RPMI 1640 medium containing 2.5% (v/v) bovine serum albumin (BSA), 0.2% (v/v) sodium azide and 20 mM Hepes, pH 7.4 binding medium. The final pool of ¹²⁵I CD40/Fc was diluted to a working stock solution of 1 x 10-7 M in binding medium and stored for up to one month at 4° C without detectable loss of receptor binding activity.

Approximately 50% - 60% label incorporation was observed. Radioiodination yielded specific activities in the range of 1×10^{15} to 5×10^{15} cpm/nmole (0.42 - 2.0 atoms of radioactive iodine per molecule of protein). SDS polyacrylamide gel electrophoresis (SDS-PAGE) revealed a single labeled polypeptide consistent with expected values. The labeled fusion protein was greater than 98% trichloroacetic acid (TCA) precipitable, indicating that the 125 I was covalently bound to the protein.

EXAMPLE 2

This example describes selection of a cell line putatively expressing CD40-L. Several cell lines were screened using the radioiodinated CD40/Fc fusion protein described in Example 1. Briefly, quantitative binding studies were performed according to standard methodology, and Scatchard plots were derived for the various cell lines. A clonal cell line (EL4, ATCC Catalog TIP 39) a murine thymoma cell line was identified and sorted. Prior to sorting, EL-4 cells were found to express approximately 450 molecules of CD40-L per cell. The seventh sort cells were called EL-40.7 and were grown and found to express approximately 10,000 molecules of CD40-L per cell. Lastly, the ninth sort cells were called EL-40.9 and were grown and found to express approximately 15,000 molecules of CD40-L per cell.

EXAMPLE 3

This example describes preparation of a cDNA library for expression cloning of murine CD40-L. The library was prepared from a fifth sorted clone of a mouse thymoma cell line EL-4 (ATCC TIB 39), called EL-40.5. EL-40.5 cells were EL4 cells 5 sorted five times with biotinylated CD40/Fc fusion protein in a FACS (fluorescence activated cell sorter). A cDNA library was made from RNA obtained from EL-40.5 cells essentially as described in US Patent 4,968,607, the disclosure of which is incorporated by reference herein. Briefly, a cDNA library was constructed by reverse transcription of poly (A)+ mRNA isolated from the total RNA extracted from the EL-10 40.5 cell line. The library construction technique was substantially similar to that described by Ausubel et al., eds., Current Protocols In Molecular Biology, Vol. 1, (1987). Poly (A)+ mRNA was isolated by oligo dT cellulose chromatography and double-stranded cDNA was made substantially as described by Gubler et al., Gene 25:263, 1983. Poly(A)+ mRNA fragments were converted to RNA-cDNA hybrids by 15 reverse transcriptase using random hexanucleotides as primers. The RNA-cDNA hybrids were then converted into double-stranded cDNA fragments using RNAase H in combination with DNA polymerase I. The resulting double-stranded cDNA was bluntended with T4 DNA polymerase.

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5'- TCG ACT GGA ACG AGA CGA CCT GCT -3'
GA CCT TGC TCT GCT GGA CGA - 5'

were ligated to 5' ends of resulting blunt-ended cDNA, as described in Haymerle et al., *Nucleic Acids Res.* 14:8615, 1986. Non-ligated adaptors were removed by gel filtration chromatography at 68°C. This left 24 nucleotide non-self-complementary overhangs on cDNA. The same procedure was used to convert 5' *Sal* I ends of the mammalian expression vector pDC406 to 24 nucleotide overhangs complementary to those added to cDNA. Optimal proportions of adaptored vector and cDNA were ligated in the presence of T4 polynucleotide kinase. Dialyzed ligation mixtures were electroporated into *E. coli* strain DH5α and transformants selected on ampicillin plates.

Plasmid DNA was isolated from pools consisting of approximately 2,000 clones of transformed *E. coli* per pool. The isolated DNA was transfected into a subconfluent layer of CV1-EBNA cells using DEAE-dextran followed by chloroquine treatment substantially according to the procedures described in Luthman et al., *Nucl. Acids Res.* 11:1295, 1983 and McCutchan et al., *J. Natl. Cancer Inst.* 41:351, 1986.

CV1-EBNA cells were maintained in complete medium (Dulbecco's modified Eagles' media containing 10% (v/v fetal calf serum, 50 U/ml penicillin, 50 U/ml streptomycin, and 2 mM L-glutamine) and were plated to a density of approximately 2

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x 10⁵ cells/well in single-well chambered slides (Lab-Tek). The slides were pre-treated with 1 ml human fibronectin (10 μg/ml PBS) for 30 minutes followed by a single washing with PBS. Media was removed from adherent cells growing in a layer and replaced with 1.5 ml complete medium containing 66.6μM chloroquine sulfate. About 0.2 ml of a DNA solution (2 μg DNA, 0.5 mg/ml DEAE-dextran in complete medium containing chloroquine) was added to the cells and the mixture was incubated at 37°C for about five hours. Following incubation, media was removed and the cells were shocked by addition of complete medium containing 10% DMSO (dimethylsulfoxide) for 2.5 - 20 minutes. Shocking was followed by replacement of the solution with fresh complete medium. The cells were grown in culture for two to three days to permit transient expression of the inserted DNA sequences. These conditions led to a 30% to 80% transfection frequency in surviving CV1-EBNA cells.

EXAMPLE 4

This example describes screening of the expression cloning library made in Example 3 with a labeled CD40/Fc fusion protein made in Example 1. After 48 - 72 hours, transfected monolayers of CV1-EBNA cells made in Example 3 were assayed by slide autoradiography for expression of CD40-L using radioiodinated CD40/Fc fusion protein as prepared in Example 1. Transfected CV1-EBNA cells were washed once with binding medium (RPMI 1640 containing 25 mg/ml bovine serum albumin (BSA), 2 mg/ml sodium azide, 20 mM Hepes pH 7.2, and 50 mg/ml nonfat dry milk) and incubated for 2 hours at 4°C ml in binding medium containing 1 x 10-9 M 125I-CD40/Fc fusion protein. After incubation, cells in the chambered slides were washed three times with binding buffer, followed by two washes with PBS, (pH 7.3) to remove unbound radiolabeled fusion protein.

The cells were fixed by incubating in 10% gluteraldehyde in PBS (30 minutes at room temperature), washed twice in PBS and air-dried. The slides were dipped in Kodak GTNB-2 photographic emulsion (6x dilution in water) and exposed in the dark for two to four days days at room temperature in a light-proof box. The slides were developed in Kodak D19 developer, rinsed in water and fixed in Agfa G433C fixer. The slides were individually examined under a microscope at 25-40x magnification. Positive slides showing cells expressing CD40-L were identified by the presence of autoradiographic silver grains against a light background.

One pool containing approximately 2000 individual clones was identified as potentially positive for binding the CD40/Fc fusion protein. The pool was titered and plated to provide plates containing approximately 200 colonies each. Each plate was scraped to provide pooled plasmid DNA for transfection into CV1-EBNA cells

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instructions.

according to the same procedure described above. The smaller pools were screened by slide autoradiography as described previously. One of the smaller pools contained clones that were positive for CD40-L as indicated by the presence of an expressed gene product capable of binding to the CD40/Fc fusion protein.

The positive smaller pool was titered and plated to obtain individual colonies. Approximately 400 individual colonies were picked and inoculated into culture medium in individual wells of 96-well plates. Cultures were mixed by pooling rows and columns and the mixed cultures were used to prepare DNA for a final round of transfection and screening. An intersection of a positive row and and a positive column indicated a potential positive colony. Ten potential positive colonies (i.e., candidate clones) were identified. DNA was isolated from each candidate clone, retransfected and rescreened. Five candidate clones were positive by binding to CD40/Fc. All five positive candidate clones contained a cDNA insert of 1468 nucleotides, as determined by dideoxynucleotide sequencing. The cDNA coding region of the CD40-L clone corresponds to the sequence of Figure 1 and SEQ ID NO:1.

A cloning vector containing murine CD40-L sequence, designated pDC406-mCD40-L, was deposited with the American Type Culture Collection, Rockville, MD (ATCC) on December 6, 1991, under accession number 68872. The nucleotide sequence and predicted amino acid sequence of this clone are illustrated in SEQ ID NO:1 and in Figure 1.

EXAMPLE 5

This example illustrates a cross-species hybridization technique which was used to isolate a human CD40-L homolog using a probe designed from the sequence of murine CD40-L. A murine CD40-L probe was produced by excising the coding region from murine CD40-L clone pDC406-CD40-L (nucleotide 13 through 793) and ³²P-labeling the fragment using random primers (Boehringer-Mannheim).

A human peripheral blood lymphocyte (PBL) cDNA library was constructed in

a λ phage vector using λgt10 arms and packaged *in vitro* using a commercially available kit (Gigapak[®] Stratagene, San Diego, CA) according to the manufacturer's instructions. The PBL cells were obtained from normal human volunteers and treated with 10 ng/ml of OKT3 (an anti-CD3 antibody), and 10 ng/ml of human IL-2 (Immunex, Seattle, WA) for six days. The PBL cells were washed and stimulated with 500 ng/ml ionomycin (Calbiochem) and 10 ng/ml PMA (Sigma) for four hours. Messenger RNA and cDNA were obtained from the stimulated PBL cells and packaged into λgt10 phage vectors (Gigapak[®] Stratagene) according to manufacturer's

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The murine probe was hybridized to phage cDNA in 6 X SSC (15 mM trisodium citrate, and 165 mM sodium chloride), 1 X Denhardt's solution, 2 mM EDTA, 0.5% Np40 at 63°C overnight. Hybridization was followed by extensive washing in 3 X SSC, 0.1% SDS at approximately 55°C for three hours. Specific bands were visualized by autoradiography.

A cloning vector containing human CD40-L sequence, designated hCD40-L, was deposited with the American Type Culture Collection, Rockville, MD (ATCC) on December 6, 1991, under accession number 68873. The nucleotide sequence and predicted amino acid sequence of this clone are illustrated SEQ ID NO:11 and in Figure 2.

EXAMPLE 6

This example illustrates the expression of membrane-bound murine CD40-L in CV1-EBNA cells. Murine CD40-L cDNA in HAVEO vector or empty HAVEO vector were transfected into CV1 EBNA cells using standard techniques, such as those described in McMahan et al. et al. *EMBO J.* 10:2821, 1991 and in Example 3 herein. Briefly, CV1 EBNA cells were plated at a density of 2 x 10⁶ cells per 10 cm dish in 10 ml of Dulbecco's Minimal Essential Medium supplemented with 10% fetal calf serum (Medium). The cells were allowed to adhere overnight at 37°C. The Medium was replaced with 1.5 ml of Medium containing 66.7 µM chloroquine and a DNA mixture containing 5 µg of cDNA encoding mCD40-L. Medium containing 175 µl, and 25 µl of DEAE dextran (4 mg/ml in PBS) was also added to the cells. The cells and cDNA were incubated at 37°C for 5 hours. The cDNA mixture was removed and the cells were shocked with 1 ml of fresh Medium containing 10% DMSO for 2.5 min. The Medium was replaced with fresh Medium and the cells were grown for at least 3 days.

EXAMPLE 7

This example illustrates the preparation of monoclonal antibodies to CD40-L. Preparations of purified murine CD40-L or human CD40-L are prepared by COS cell expression and CD40/Fc affinity purification as described herein. Purified CD40-L can generate monoclonal antibodies against CD40-L using conventional techniques, for example, those techniques described in U.S. Patent 4,411,993. Briefly, mice are immunized with CD40-L as an immunogen emulsified in complete Freund's adjuvant, and injected in amounts ranging from 10-100 µg subcutaneously or intraperitoneally. Ten to twelve days later, the immunized animals are boosted with additional CD40-L emulsified in incomplete Freund's adjuvant. Mice are periodically boosted thereafter on a weekly to bi-weekly immunization schedule. Serum samples are periodically taken

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by retro-orbital bleeding or tail-tip excision for testing by dot blot assay or ELISA (Enzyme-Linked Immunosorbent Assay), for CD40-L antibodies.

Following detection of an appropriate antibody titer, positive animals are provided one last intravenous injection of CD40-L in saline. Three to four days later, the animals are sacrificed, spleen cells harvested, and spleen cells are fused to a murine myeloma cell line (e.g., NS1 or Ag 8.653). Fusions generate hybridoma cells, which are plated in multiple microtiter plates in a HAT (hypoxanthine, aminopterin and thymidine) selective medium to inhibit proliferation of non-fused cells, myeloma hybrids, and spleen cell hybrids.

The hybridoma cells are screened by ELISA for reactivity against purified CD40-L by adaptations of the techniques disclosed in Engvall et al., *Immunochem*. 8:871, 1971 and in U.S. Patent 4,703,004. Positive hybridoma cells can be injected intraperitoneally into syngeneic BALB/c mice to produce ascites containing high concentrations of anti-CD40-L monoclonal antibodies. Alternatively, hybridoma cells can be grown *in vitro* in flasks or roller bottles by various techniques. Monoclonal antibodies produced in mouse ascites can be purified by ammonium sulfate precipitation, followed by gel exclusion chromatography. Alternatively, affinity chromatography based upon binding of antibody to protein A or protein G can also be used, as can affinity chromatography based upon binding to CD40-L.

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EXAMPLE 8

This example illustrates anti-allergy therapeutic effects of sCD40 and CD40/Fc fusion protein. Soluble CD40 and CD40/Fc were tested for their ability to inhibit IL-4 (5 ng/ml) induced IgE secretion in a two donor MLC system. The data from three experiments are presented in Table 1.

Table 1

		IgE (ng/ml)			
	Addition	Exp. 1	Exp. 2		
	Exp. 3				
30	medium	<0.1	<0.1		
	<0.1				
	IL-4	24	47	54	
	IL-4 + sCD40 (0.1 μ g/ml)	19	nd	38	
35	IL-4 + sCD40 (0.3 μ g/ml)	14	29	24	
	IL-4 + sCD40 (1 μg/ml)	10	24	8	
	IL-4 + sCD40 (3 μg/ml)	7	19	2	
	IL-4 + IL-7R/Fc (10 μg/ml)	21	nd	58	

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IgE levels were measured after 12 days in culture by an ELISA procedure. Briefly, flat-bottomed 96-well microtiter plates (Corning) were coated with mouse mAb anti-human IgE (Zymed) at 1:500 dilution in PBS (phosphate buffered saline). After washing 3X, a blocking step was performed using 5% non-fat dried milk, followed by titration of human IgE standards or test supernatants. After washing 3X, biotinylated goat anti-human IgE (Kirkegaard and Perry) was added at a 1:500 dilution. This was followed by further washing and then addition of streptavidin-HRP (Zymed) at a 1:500 dilution. After further washing, the reaction was developed using TMB substrate (Kirkegaard and Perry) and absorbance measured at 520 nm. All washing steps were carried out in PBS plus 0.05% Tween. All incubation steps were performed at volumes of 100 μl/well for one hour at room temperature. The sensitivity of this assay is 100 pg/ml.

EXAMPLE 9

This example illustrates the effects of sCD40 and CD40/Fc fusion protein to inhibit soluble CD23 shedding from IL-4 (5 ng/ml) stimulated B cells. Soluble CD40 and CD40/Fc were tested for their ability to inhibit IL-4-induced sCD23 shedding in a two donor MLC system The data from three experiments are presented in Table 2.

Table 2

20	sCD23 (ng/ml)							
	Addition	Exp. 1			Exp. 2		Exp. 3	
		day 6	day 12		day 6	day 12	day 6	
	12					•	•	•
	E- + medium	55	< 0.5		24	10	10	5
25	+ IL-4	115	55		96	62	44	27
	$+ IL-4 + sCD40 (1 \mu g/ml)$	nd	nd		88	36	38	9
	$+ IL-4 + sCD40 (3 \mu g/ml)$	97	4		82	31	40	4
	$+ IL-4 + sCD40 (10 \mu g/ml)$	nd	nd		72	28	nd	nd
	+ IL-4 + IL-7R/Fc (3 μg/ml)	111	48		103	67	40	22
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	PBM + medium	12	< 0.5		15	5	3	10
	+ IL-4	39	255		47	22	48	26
	+ IL-4 + sCD40 (1 μg/ml)	nd	nd		44	18	46	18
	+ IL-4 + sCD40 (3 μg/ml)	24	6	,	37	11	45	12
35	$+ IL-4 + sCD40 (10 \mu g/ml)$	nd	nd	:	28	5	nd	nd
	+ IL-4 + IL-7R/Fc (3 μg/ml)	35	26		43	20	50	23

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Soluble CD23 levels were measured after 6 and 12 days in culture by a commercial sCD23 ELISA detection kit (Binding Site, San Diego, CA). The sensitivity limit was 500 pg/ml. Approximately 1 x 10⁵ cells per well were cultured in triplicate in round-bottomed 96-well microtiter plates (Intermountain Scientific, Bountiful UT) for the indicated time in the presence or absence of additives as indicated in Table 2. The results show anti-allergy effects of sCD40. Similar studies were run with CD40/Fc (data not shown) instead of sCD40, and similar results were obtained. Accordingly, these data in Examples 8 and 9 illustrate an anti-allergy property for CD40.

10 EXAMPLE 10

This example illustrates B cell proliferative activity of membrane-bound murine CD40-L for human B cells. Human peripheral blood mononuclear cells (PBMC) were isolated from peripheral blood from normal volunteers by density gradient centrifugation over Histopaque® (Sigma, St. Louis, MO) T cell-depleted preparations of cells (E⁻) were obtained by removing T cells by rosetting with 2aminoethylisothiouronium bromide-treated SRBC (sheep red blood cells) and further density gradient centrifugation over Histopaque[®]. B cell proliferation assays were conducted with E- preparations in RPMI media with added 10% heat-inactivated fetal bovine serum (FBS) at 37°C in a 10% CO₂ atmosphere. Approximately 1 X 10⁵ E⁻ cells per well were cultured in triplicate in flat-bottomed 96-well microtiter plates (Corning) for 7 days in the presence of transfected CV1 EBNA cells (described in Example 6). The CV1 EBNA cells were transfected with murine CD40-L cDNA or empty vector. The cells were pulsed with 1 µCi/well of tritiated thymidine (25 Ci/nmole Amersham, Arlington Heights, IL) for the final eight hours of culture. Cells were harvested onto glass fiber discs with an automated cell harvester and incorporated cpm were measured by liquid scintillation spectrometry.

Figure 4a shows a comparison of human B cell proliferation of CV1 EBNA cells transfected with empty vector (HAVEO) or with murine CD40-L cDNA in HAVEO vector. These data show that membrane-bound CD40-L stimulates human B cell proliferation in the absence of a co-mitogen. Figure 4b shows a similar experiment, except that 10 ng/ml of human IL-4 was added to the cultures. In this experiment, IL-4 slightly enhances the B cell mitogenic activity of membrane-bound murine CD40-L. Figure 5 is a repeat of the experiment shown in Figure 4b. However, when the experiment was repeated, there was no evidence of IL-4 co-mitogenic activity. There was repeated evidence of CD40-L mitogenic activity. Accordingly, membrane-bound CD40-L stimulates proliferation of human B cells.

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35 EXAMPLE 11

This example illustrates the effect of membrane-bound murine CD40-L to stimulate IgE production and CD23 shedding from E⁻ cells isolated in Example 10. Approximately 1 X 10⁵ cells/well were cultured in triplicate round bottomed 96-well Nunc microtiter plates (Intermountain Scientific, Bountiful UT) in Iscove's Modified Dulbecco's Medium (IMDM) plus 10% FCS in a humidified atmosphere of 10% CO₂. Medium was supplemented with 50 μg/ml human transferrin (Sigma), 0.5% bovine serum albumin (Sigma) and 1 μg/ml of each of oleic, linoleic and palmitic acids (Sigma). The E⁻ cells were cultured for 10 days in the presence of 5 ng/ml human IL-10 4. A titration of CV1 EBNA cells transfected with murine CD40-L or empty vector were added. After ten days, culture supernatants were assayed for IgE by the ELISA procedure described in Example 8 or for CD23 shedding by the procedure described in Example 9.

Figure 6 shows a comparison of IgE production in the supernatants (in ng/ml) for cultures of E- cells and CV1 EBNA cells transfected with empty vector (HAVEO) 15 or with CD40-L. No differences were noted with up to 3000 CV1 EBNA cells, however significant IgE production resulted with the addition of 10000 or 30000 CD40-L transfected CV1 EBNA cells. As a comparison, when E- cells were incubated with medium alone, 5 ng/ml IL-4 or 5 ng/ml IL-4 plus 500 ng/ml G28-5 antibody, IgE 20 production was 4.7, 2.9 and >600 ng/ml, respectively. When CD23 shedding was measured in Figure 7, 10000 and 30000 CVI EBNA cells transfected with CD40-L showed increased CD23 shedding when compared to empty vector control CV1 EBNA cells. As a comparison, when E- cells were incubated with medium alone, 5 ng/ml IL-4 or 5 ng/ml IL-4 plus 500 ng/ml G28-5 antibody, CD23 shedding was <0.1, 2.4 and 25 11.2 ng/ml, respectively. These data show that IgE production and CD23 shedding are both biological activities associated with membrane-bound CD40-L.

EXAMPLE 12

This example illustrates B cell proliferative activity, polyclonal immunoglobulin

(Ig) production, antigen-specific antibody formation and various method for using membrane-bound and soluble CD40-L in clinical applications. We obtained murine splenic B cells according to procedures described in Grabstein et al. I supra, Maliszewski et al. I supra and Maliszewski et al. II supra. Briefly, the mixed culture of cells was purified by T cell depletion using T cell antiserum and complement, and adherent cell depletion by passage of Sephadex® G10 columns and by B cell positive selection by panning on petri dishes coated with goat anti-mouse IgM. Purified B cells were cultured in RPMI, fetal calf serum (5% for B cell proliferation assays and 20% for

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plaque forming cell assays or polyclonal antibody assays), 2-mercaptoethanol, antibiotics, amino acids and pyruvate. B cell proliferation was measured according to the assay described in Example 10 and in Grabstein et al. I supra, Maliszewski et al. I supra and Maliszewski et al. II supra. Antigen-specific antibody formation was

5 measured by the procedure described in Grabstein et al., J. Mol. Cell. Immunol. 2:199, 1986 [Grabstein et al. II]. Briefly, antigen specific antibody formation used sheep red blood cells (SRBC) as antigen (0.03% v/v) in 2.0 ml cultures of 1 X 106 murine B cells per culture. The B cell cultures were incubated for 5 days and plaque forming cells were determined by Jerne hemolytic plaque assay as described in Grabstein et al.

10 II supra. Cell counts were determined in a coulter counter. Polyclonal Ig secretion was determined by isotype-specific ELISA assays in seven day cultures of 1 X 106 B cells per 2.0 ml culture as described in Maliszewski et al. I supra and Maliszewski et al. II supra.

The results of B cell proliferation by CV1 EBNA cells transfected with CD40-L or empty vector or 7A1 cells (a T cell helper clone) are shown in Figures 8, 10 and 12. These data show that the greatest B cell proliferation was caused by CD40-L. T cell helper cells 7A1 and 7C2 had a minimal effect on B cell proliferation.

The effects of various cells upon antigen specific antibody formation are shown in Figures 9 and 11. Figure 9 shows a comparison of plaque forming cells comparing T cell helper clone 7A1 and murine EL40.9 cells which secrete a soluble CD40-L. The EL40.9 cells seem to have an inhibitory effect upon antigen specific antibody formation. Figure 11 shows PFC (plaque forming cells) for T cell helper cells 7C2 and CV1 EBNA cells transfected with either empty vector or CD40-L. Both 7C2 cells and membrane-bound CD40-L stimulated antigen specific antibody formation (PFC).

Figure 13 compares antigen specific antibody formation of CD40-L and 7A1 cells in the

Figure 13 compares antigen specific antibody formation of CD40-L and 7A1 cells in the presence or absence of 10 ng/ml interleukin-2 (IL-2). IL-2 increased PFC for 7A1 cells but did not increase PFC caused by membrane-bound CD40-L.

Polyclonal Ig production by murine B cells was compared for stimulation or inhibition with membrane-bound CD40-L, control CV1 EBNA cells and helper T cells 7A1 in the presence of cytokines IL-4 (10 ng/ml) and IL-5 (1:40 dilution of COS cell supernatants) or without added cytokines. The amount of IgA, IgG3, IgE, IgG2b, IgM and IgG1 are shown in Tables 3-8, respectively.

TABLE 3

IgA, ng/ml

			-	-
5		# CELLS	MEDIA	+IL-4+IL-5
10	CD40-L	2 X 10(5) 1 X 10(5) 1 X 10(4)	666.275 ± 174.444 288.085 ± 20.773 53.750 ± 36.531	64.639 ± 51.780 291.831 ± 10.673 910.072 ± 62.713
15	HAVEO	2 X 10(5) 1 X 10(5) 1 X 10(4)	0 0 0	628.190 ± 42.907 477.755 ± 57.478 295.640 ± 12.736
13	7A1 (2C11)	1 X 10(6) 2 X 10(5) 1 X 10(5)	0 0 0	2177.549 ± 377.052 646.898 ± 86.325 480.671 ± 40.011
20	MEDIA LPS		$ \begin{array}{c} 0 \\ 88.531 \pm 31.248 \end{array} $	458.152 ± 77.258 132.336 ± 51.356
25			TABLE 4	
			IgG3, ng/ml	
30		# CELLS	MEDIA	+IL-4+IL-5
35	CD40-L	2 X 10(5) 1 X 10(5) 1 X 10(4)	108.427 ± 14.359 118.079 ± 8.021 127.591 ± 6.268	0 46.535 ± 9.899 467.023 ± 78.276
40	HAVEO	2 X 10(5) 1 X 10(5) 1 X 10(4)	$0 \\ 11.205 \pm 4.434 \\ 26.389 \pm 10.221$	29.773 ± 5.224 66.323 ± 8.673 34.671 ± 12.975
.5	7A1 (2C11)	1 X 10(6) 2 X 10(5) 1 X 10(5)	33.420 ± 9.972 0 0	820.856 ± 39.442 436.074 ± 59.332 239.760 ± 45.978
45	MEDIA LPS		21.808 ± 7.107 816.697 ± 43.553	64.773 ± 13.924 103.720 ± 11.883

38 • TABLE 5

IgE, ng/ml

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3		# CELLS	MEDIA	+IL-4+IL-5
10	CD40-L	2 X 10(5) 1 X 10(5) 1 X 10(4)	0 0 0	64.144 ± 4.979 83.493 ± 9.093 461.155 ± 60.514
15	HAVEO	2 X 10(5) 1 X 10(5) 1 X 10(4)	0 0 0	$\begin{array}{c} 0 \\ 4.208 \pm .527 \\ 0 \end{array}$
13	7A1 (2C11)	1 X 10(6) 2 X 10(5) 1 X 10(5)	0 0 0	208.091 ± 8.090 32.530 ± 0.723 15.889 ± 2.947
20	MEDIA LPS		0 0	12.602 ± 1.460 408.355 ± 9.764
25			TABLE 6	
			IgG2b, ng/ml	
30		# CELLS	MEDIA	+IL-4+IL-5
35	CD40-L	2 X 10(5) 1 X 10(5) 1 X 10(4)	0 0 0	0 6.230 ± .285 47.414 ± .241
40	HAVEO	2 X 10(5) 1 X 10(5) 1 X 10(4)	0 0 0	7.001 ± 2.358 6.230 ± 2.285 9.620 ± 2.650
4 0	7A1 (2C11)	1 X 10(6) 2 X 10(5) 1 X 10(5)	0 0 0	189.343 ± 2.837 22.431 ± 6.835 7.207 ± 1.580
45	MEDIA LPS		0	7.422 ± 1.602 33.291 ± 3.183

39 TABLE 7

IgM, $\mu g/ml$

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		# CELLS	MEDIA	+IL-4+IL-5
10	CD40-L	2 X 10(5) 1 X 10(5) 1 X 10(4)	1.805 ± 0.639 2.237 ± 0.583 2.293 ± 0.595	0.439 ± 0.184 5.878 ± 0.858 96.730 ± 13.009
15	HAVEO	2 X 10(5) 1 X 10(5) 1 X 10(4)	$0 \\ 0 \\ 0.624 \pm 0.178$	10.890 ± 2.126 13.303 ± 0.993 22.538 ± 2.304
20	7A1 (2C11)	1 X 10(6) 2 X 10(5) 1 X 10(5)	0.769 ± 0.124 0.142 ± 0.052 0.126 ± 0.048	104.857 ± 17.463 27.016 ± 1.706 13.070 ± 0.600
20	MEDIA LPS		0.231 ± 0.057 53.302 ± 9.668	36.809 ± 2.860 41.974 ± 6.158
25			TABLE 8	
			IgG1, ng/ml	
30		# CELLS	MEDIA	+IL-4+IL-5
	CD40-L	2 X 10(5) 1 X 10(5) 1 X 10(4)	0 0 0	130.185 ± 24.547 310.588 ± 1.261
35		1 7 10(4)	U	270.727 ± 17.511
40	HAVEO	2 X 10(5) 1 X 10(5) 1 X 10(4)	0 0 0	187.668 ± 57.730 43.320 ± 49.770 1363.464 ± 45.841
	7A1 (2C11)	1 X 10(6) 2 X 10(5) 1 X 10(5)	0 0 0	145.652 ± 136.070 365.563 ± 24.276 449.475 ± 101.012
45	MEDIA LPS		0	133.660 ± 386.231 246.213 ± 21.526

These data indicate that the interaction of CD40 with its ligand is the principal molecular interaction responsible for T cell contact dependent induction of B cell growth and differentiation to both antigen-specific antibody production and polyclonal Ig secretion. As such, these data suggest that antagonists of this interaction, by soluble CD40, CD40/Fc fusion protein and possibly soluble CD40-L (monomeric), will

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significantly interfere with development of antibody responses. Therefore clinical situations where CD40, CD40/Fc fusion proteins and soluble CD40-L include allergy, lupus, rheumatoid arthritis, insulin dependent diabetes mellitus, and any other diseases where autoimmune antibody or antigen/antibody complexes are responsible for clinical pathology of the disease. Moreover, membrane-bound CD40-L or oligomeric soluble CD40-L will be useful to stimulate B cell proliferation and antibody production. As such, these forms of CD40-L are most useful for vaccine adjuvants and as a stimulating agent for mAb secretion from hybridoma cells.

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EXAMPLE_13

This example illustrates the effect of membrane-bound CD40-L upon proliferation of and IgE secretion from peripheral blood mononuclear cells (E⁻). Ecells were obtained according to the procedure described in Example 10 and incubated for 7 or 10 days in the presence of CV1 EBNA cells transfected with empty vector or mCD40-L cDNA. Additionally, CD40/Fc fusion protein (described in Example 1) or TNF Receptor/Fc fusion protein (described in WO 91/03553) was added to some of the preparations as indicated in Figure 14. IgE secretion was measured according to the procedure described in Example 8 and B cell proliferation was measured according to the procedure described in Example 10.

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The results for B cell proliferation and IgE secretion are shown in Figure 14 for five different concentrations of transfected CV1 EBNA cells. Both B cell proliferation and IgE secretion were increased in the presence of membrane-bound CD40-L. Addition of CD40/Fc fusion protein ablated both B cell proliferation and IgE secretion. The TNF Receptor/Fc fusion protein had no effect. As a comparison for IgE secretion, addition of IL-4 as a control agent (without transfected CV1 EBNA cells) produced no IgE in this assay and addition of IL-4 plus G28-5 anti-CD40 mAb resulted in 29.7 ng/ml IgE in this assay.

EXAMPLE 14

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This example describes construction of a CD40-L/Fc DNA construct to express a soluble CD40-L/Fc fusion protein referred to as CD40-L/FC2 construct. DNA encoding CD40-L/FC2 comprises sequences encoding a leader (or signal) peptide, an eight amino acid hydrophilic sequence described by Hopp et al. (Hopp et al., *Bio/Technology* 6:1204,1988; referred to as Flag[®]), a suitable Fc region of an immunoglobulin, a [Gly4Ser]₃ repeat sequence (described in U.S. Patent 5,073,627, which is incorporated by reference herein) or other suitable linker sequence, and the extracellular region of human CD40-L from amino acid 50 to amino acid 261 (SEQ ID

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NO:11). A pDC406 expression vector containing a leader sequence, Flag[®], and human IgG₁ Fc is prepared using conventional techniques of enzyme cutting and ligation of fragments encoding a leader sequence, Flag[®], and human IgG₁ Fc, and restricted with *Nsi* 1 and *Not* 1.

A PCR technique (Sarki et al., Science 239:487, 1988) was employed using 5' (upstream) and 3' (downstream) oligonucleotide primers to amplify the DNA sequences encoding CD40 extracellular ligand binding domain from a cloning vector containing human CD40-L (ATCC 68873; SEQ ID NO: 11) to form a PCR fragment. The upstream oligonucleotide primer (SEQ ID NO:13) introduced a Nsi 1 site upstream from a linker sequence ([Gly4Ser]3SerSer), which was followed by 21 nucleotides of the extracellular domain of CD40-L (amino acids 51 through 57 of SEQ ID NO: 11). A downstream oligonucleotide primer (SEQ ID NO:14) introduced a Not 1 site just downstream of the termination codon of the CD40-L. The PCR fragment was then ligated into the pDC406 expression vector containing a leader sequence, Flag®, and human IgG₁ Fc. The nucleotide and predicted amino acid sequence of CD40-L/FC2 are presented in SEQ ID NO:15 and SEQ ID NO:16. The resultant DNA construct (CD40-L/FC2) was transfected into the monkey kidney cell line CV-1/EBNA (ATCC CRL 10478). The construct encoded a soluble CD40-L capable of binding CD40, as evidenced by binding observed in fluorescence-activated cell sorting (FACS) analysis using cells that express CD40.

Large scale cultures of human embryonic kidney 293 cells (ATCC CRL 1573) transfected with the construct encoding CD40-L/FC2 were grown to accumulate supernatant containing CD40-L/FC2. The 293 cell line, a permanent line of primary human embryonal kidney transformed by human adenovirus 5 DNA, permits expression of recombinant proteins ligated into the pCD406 vector. The CD40-L/FC2 fusion protein in supernatant fluid was purified by affinity purification. Briefly, culture supernatant containing the CD40-L/FC2 fusion protein was purified by filtering mammalian cell supernatants (e.g., in a 0.45μ filter) and applying filtrate to an antibody affinity column comprising biotinylated goat anti-human IgG (Jackson Immunoresearch Laboratories, Inc., Westgrove, PA, USA) coupled to Streptavidin-agarose (Pierce Chemical, Rockford, IL, USA) at 4°C, at a flow rate of approximately 60 to 80 ml/hr for a 1.5 cm x 12.0 cm column. The column was washed with approximately 20 column volumes of PBS (phosphate buffered saline), until free protein could not be detected in wash buffer. Bound fusion protein was eluted from the column with 12.5 mM citrate buffer, 75 mM NaCl, pH 2.8, and brought to pH 7 with 500 mM Hepes buffer, pH 9.1. The purified, oligomeric CD40-L/FC2 peptide induced human B cell proliferation in the absence of any co-stimuli, and (in conjunction with the appropriate

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cytokine) resulted in the production of IgG, IgE, IgA and IgM, as described in Example 12 for membrane-bound CD40-L.

EXAMPLE 15

This example describes construction of a CD40-L DNA construct to express a soluble CD40-L fusion protein referred to as trimeric CD40-L. Trimeric CD40-L contains a leader sequence, a 33 amino acid sequence referred to as a "leucine zipper" (SEQ ID NO:17), and an eight amino acid hydrophilic sequence described by Hopp et al. (Hopp et al., *Bio/Technology* 6:1204,1988; referred to as Flag[®]), followed by the extracellular region of human CD40-L from amino acid 50 to amino acid 261 (SEQ ID NO:11). The utility of the leader and the Flag[®] sequences have been described in the Detailed Description. The 33 amino acid sequence presented in SEQ ID NO:17 trimerizes spontaneously in solution. Fusion proteins comprising this 33 amino acid sequence are thus expected to form trimers or multimers spontaneously.

The construct is prepared by synthesizing oligonucleotides representing a leader sequence, the 33 amino acid sequence described above, and the Flag[®] sequence, then ligating the final product to a DNA fragment encoding amino acids 51 through 261 of SEQ ID NO:11, prepared as described in Example 14.

The resulting ligation product in expression vector pDC406 was transfected into the monkey kidney cell line CV-1/EBNA (ATCC CRL 10478). The pDC406 plasmid includes regulatory sequences derived from SV40, human immunodeficiency virus (HIV), and Epstein-Barr virus (EBV). The CV-1/EBNA cell line was derived by transfection of the CV-1 cell line with a gene encoding Epstein-Barr virus nuclear antigen-1 (EBNA-1) that constitutively expresses EBNA-1 driven from the human CMV intermediate-early enhancer/promoter. The EBNA-1 gene allows for episomal replication of expression vectors, such as pDC406, that contain the EBV origin of replication.

Once cells expressing the fusion construct are identified, large scale cultures of transfected cells are grown to accumulate supernatant from cells expressing trimeric CD40-L. The trimeric CD40-L fusion protein in supernatant fluid is purified by affinity purification substantially as described in U.S. Patent 5,011,912. Silver-stained SDS gels of the eluted CD40-L fusion protein can be prepared to determine purity.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

40

(i) APPLICANT: ARMITAGE, RICHARD

FANSLOW, WILLIAM SPRIGGS, MELANIE

5	(ii) TITLE OF INVENTION: NOVEL CYTOKINE
3	(iii) NUMBER OF SEQUENCES: 17
10	(iv) CORRESPONDENCE ADDRESS: (A) ADDRESSEE: IMMUNEX CORPORATION (B) STREET: 51 UNIVERSITY STREET (C) CITY: SEATTLE (D) STATE: WASHINGTON (E) COUNTRY: USA (F) ZIP: 98101
15	(v) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk
20	(B) COMPUTER: IBM PC compatible(C) OPERATING SYSTEM: PC-DOS/MS-DOS(D) SOFTWARE: PatentIn Release #1.0, Version #1.25
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25	(B) FILING DATE: (C) CLASSIFICATION:
	(viii) ATTORNEY/AGENT INFORMATION:
	(A) NAME: OSTER, JEFFREY B. (B) REGISTRATION NUMBER: 32585
30	(C) REFERENCE/DOCKET NUMBER: 2802
35	(ix) TELECOMMUNICATION INFORMATION: (A) TELEPHONE: 2065870430 (B) TELEFAX: 2065870606
	(2) INFORMATION FOR SEQ ID NO:1:
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 783 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
45	(ii) MOLECULE TYPE: cDNA
	(iii) HYPOTHETICAL: NO
50	(iv) ANTI-SENSE: NO
	(vi) ORIGINAL SOURCE: (A) ORGANISM: MOUSE
55	(vii) IMMEDIATE SOURCE: (B) CLONE: CD40-L
60	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1783

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

	A 4		ATA	A GZ	AA A	CA 1	AC	AGO	CA	A C	CT	TCC	CC	C A	GA :	rcc	GT	G G	CA A	CT.	GGA
5		-	Ile	e GI	Lu T	hr I	'y <i>‡</i> 5	Ser	G1	n P	ro	Ser		:0 A .0	rg S	Ser	۷a	1 A		hr 15	Gly
	C:		CCA	GC	CG A	GC A	TG.	AAG	AT	T T	ГT	ATG	TA	T T:	ra c	TT	AC:	r G1	T T	TC	CTT
10			Pro	Al		er M 20	et	Lys	Il	e Pl	ne	Met 25		r Le	eu I	eu	Th	_	1 P	he	Leu
	A7 14		ACC	CA	A A	G A	TT	GGA	TC	T GI	rG	CTT	TT	T GO	CT G	TG	TAT	CI	T C	ΑT	AGA
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	AG 19		TG	GA	T AA	G G	rc (GAA	GA	G GA	A (GTA	AA	CI	T C	ΑT	GAA	GA	T TI	T	GTA
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25		e I	le	Lys	s Ly	s Le	eu 1	Lys 70	Arg	ГСУ	s F	Asn	Lys	Gl _.	_	Lu	Gly	Se	r Le	u :	Ser 80
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40			g (Gly 115	Asp	Glı	ı A	sp 1	Pro	Gln 120		le i	Ala	Ala	Hi.		7al .25	Val	Ser	G	lu
	GCC 432		.C I	AGT	AAT	GCZ	A GO	CA :	rcc	GTT	C	CA (CAG	TGG	GC	C A	AG	AAA	GGA	T	AT
45				Ser	Asn	Ala	a Al		Ser 135	Val	Le	eu (ln	Trp	Ala 140		ys	Lys	Gly	T	yr
	TAT 480	AC	C A	TG	AAA	AGC	: A2	AC I	TG	GTA	AT	G C	TT	GAA	AAT	G	GG 2	AAA	CAG	C?	rg
50		Th	r M	let	Lys	Ser	15		eu	Val	Ме	t I	eu	Glu 155	Asn	ı G.	ly :	Lys	Gln		∋u 50
	ACG 528	GT:	r a	AA	AGA	GAA	GG	A C	TC	TAT	TA	T G	TC	TAC	ACI	. Cz	AA (STC	ACC	TI	rc
55	Thr	Va]	L L	ys	Arg	Glu 165	Gl	y L	eu	Tyr	Ту		al 70	Tyr	Thr	G.	ln V	/al	Thr 175	Ph	ıe
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60	Cys	Ser	: A:	sn .	Arg	Glu	Pr	o S	er :	Ser	Gl:	n. A:	rg 1	?ro	Phe	IJ		al	Gly	Le	u

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4			Leu	Lys 195	Pro	Se	r S	er (Gly	Se 20	r G O	lu	Arg	Ile	Let		eu 05	Lys	Ala	a Ala	
	A 6	AT . 72	ACC	CAC	AGT	TC	C T	CC (CAG	CT	r T	GC	GAG	CAG	CAC	T (CT	GTT	CAC	TTG	
10	A	sn	Thr 210	His	Ser	Se:	r Se	er (31n 215	Let	1 C	ys	Glu	Gln	Glr 220	ı Se	er '	Val	His	Leu	
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					(B) (D)		PE: POLC														
35			(ii	.) MC	LECU	JLE	TYP	E:	pro	tei	n										
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45	Ile	Th	r G	ln M	et I	le	Gly	Sei	r Va	al 1 40	Leu	Ph	e Al	la V	al 1	'yr 45	Le	u H	is A	Arg	
50	Arg	Le:	u As O	sp Ly	ys V	al (Glu	Gl t 55	1 G.	lu V	/al	As	n Le	eu H	is G 60	Slu	As	p Pl	he V	/al	
	Phe 65	Ile	e Ly	s Ly	s Le	eu 1	Lys 70	Arg	ı Cž	/S A	sn	Lу	s Gl 7	y G. '5	lu G	ly	Se	r Le	eu S	er 80	
55	Leu	Let	ı As	n Cy	rs G]	lu 6 35	Glu	Met	Ar	g A	.rg	Gli 90	n Ph O	e G	u A	sp	Leı		al L	ys	
	Asp	Ile	Th	r Le 10	ú As O	sn I	ys	Glu	Gl	u L	ys 05	Lys	s Gl	u As	sn S		Phe 110		u M	et	
60	Gln	Arg	Gl:	y As	p Gl	u A	sp :	Pro	Gl 12	n I.	le.	Ala	a Al	a Hi	s Va	al '	Val	. Se	r G	lu	

	Ala Asn 130		Asn	Ala	Ala	135		Leu	Gln	Trp	A1a 140	Lys	тÀз	GTA	Tyr
5	Tyr Thr 145	Met	Lys	Ser	Asn 150	Leu	Val	Met	Leu	Glu 155	Asn	Gly	Lys	Gln	Leu 160
10	Thr Val	Lys	Arg	Glu 165		Leu	Tyr	Tyr	Val 170	Tyr	Thr	Gln	Val	Thr 175	Phe
10	Cys Ser		Arg 180	Glu	Pro	Ser	Ser	Gln 185	Arg	Pro	Phe	Ile	Val 190	Gly	Leu
15	Trp Leu	Lys 195	Pro	Ser	Ser	Gly	Ser 200	Glu	Arg	Ile	Leu	Leu 205	Lys	Ala	Ala
	Asn Thr 210	His	Ser	Ser	Ser	Gln 215	Leu	Cys	Glu	Gln	Gln 220	Ser	Val	His	Leu
20	Gly Gly 225	Val 1	Phe		Leu 230	Gln	Ala	Gly		Ser 235	Val	Phe	Val	Asn	Val 240
25	Thr Glu	Ala :		Gln 245	Val	Ile	His	_	Val 250	Gly	Phe	Ser	Ser	Phe 255	Gly
	Leu Leu	-	Leu 260												
30	(2) INFO	RMATI	ON 1	FOR S	SEQ :	ID N	0:3:								
35	(i)	(B) (C)	LEN TYE STE	E CHA NGTH: PE: 1 RANDE	: 740 nucle EDNES	0 bas eic a SS: s	se pacid	airs							
40		MOLE													
40	(iii) (iv)	HYPO ANTI)									
45	(vi)			SOU ANIS			Г								
	(vii)			E SO		-									
50	(xi)	SEQUE	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	3:					
55	CGGTACCGC	T AGO	GTC	GACA	GGC	CTAG	GAT .	ATCG	ATAC	GT A	GAGC	CCAG	A TC	TTGT	GACA
JJ	AAACTCACA	C ATG	CCCZ	ACCG	TGC	CCAG	CAC (CTGA	ACTC	CT GO	GGGG	GACC	G TC	AGTC:	TTCC
60	TCTTCCCCC	C AAA	ACCO	CAAG	GAC	ACCCI	CA 1	GATO	CTCCC	CG GZ	vccc.	CTGA	GT	CACA	rgcg

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	TGGTGGTGGA CGTGAGCCAC GAAGACCCTG AGGTCAAGTT CAACTGGTAC GTGGACGGCG 240
5	TGGAGGTGCA TAATGCCAAG ACAAAGCCGC GGGAGGAGCA GTACAACAGC ACGTACCGGG
	TGGTCAGCGT CCTCACCGTC CTGCACCAGG ACTGGCTGAA TGGCAAGGAC TACAAGTGCA
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15	AGCCCCGAGA ACCACAGGTG TACACCCTGC CCCCATCCCG GGATGAGCTG ACCAAGAACC
13	AGGTCAGCCT GACCTGCCTG GTCAAAGGCT TCTATCCCAG GCACATCGCC GTGGAGTGGG 540
20	AGAGCAATGG GCAGCCGGAG AACAACTACA AGACCACGCC TCCCGTGCTG GACTCCGACG
	GCTCCTTCTT CCTCTACAGC AAGCTCACCG TGGACAAGAG CAGGTGGCAG CAGGGGAACG
25	TCTTCTCATG CTCCGTGATG CATGAGGCTC TGCACAACCA CTACACGCAG AAGAGCCTCT 720
30	CCCTGTCTCC GGGTAAATGA 740
	(2) INFORMATION FOR SEQ ID NO:4:
35	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 519 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
40	(ii) MOLECULE TYPE: cDNA
	(iii) HYPOTHETICAL: NO
•	(iv) ANTI-SENSE: NO
45	(vi) ORIGINAL SOURCE: (A) ORGANISM: HUMAN
50	(vii) IMMEDIATE SOURCE: (B) CLONE: CD40 EXTRACELLULAR REGION
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
55	CAGAACCACC CACTGCATGC AGAGAAAAAC AGTACCTAAT AAACAGTCAG TGCTGTTCTT
	TGTGCCAGCC AGGACAGAAA CTGGTGAGTG ACTGCACAGA GTTCACTGAA ACGGAATGCC 120
60	TTCCTTGCGG TGAAAGCGAA TTCCTAGACA CCTGGAACAG AGAGACACAC TGCCACCAGC 180

	ACAAATACTG CGACCCCAAC CTAGGGCTTC GGGTCCAGCA GAAGGGCACC TCAGAAACAG 240
5	ACACCATCTG CACCTGTGAA GAAGGCTGGC ACTGTACGAG TGAGGCCTGT GAGAGCTGTG
10	TCCTGCACCG CTCATGCTCG CCCGGCTTTG GGGTCAAGCA GATTGCTACA GGGGTTTCTG 360
10	ATACCATCTG CGAGCCCTGC CCAGTCGGCT TCTTCTCCAA TGTGTCATCT GCTTTCGAAA 420
15	AATGTCACCC TTGGACAAGC TGTGAGACCA AAGACCTGGT TGTGCAACAG GCAGGCACAA 480
	ACAAGACTGA TGTTGTCTGT GGTCCCCAGG ATCGGCTGA 519
20	(2) INFORMATION FOR SEQ ID NO:5:
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: cDNA
30	(iii) HYPOTHETICAL: NO
	(iv) ANTI-SENSE: NO
35	(vi) ORIGINAL SOURCE: (A) ORGANISM: PCR PRIMER
	(vii) IMMEDIATE SOURCE: (B) CLONE: CD40 5' PRIMER
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:
45	CCGTCGACCA CCATGGTTCG TCTGCC 26
	(2) INFORMATION FOR SEQ ID NO:6:
50	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 28 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
55	(ii) MOLECULE TYPE: cDNA
	(iii) HYPOTHETICAL: NO
	(iv) ANTI-SENSE: NO
60	(vi) ORIGINAL SOURCE: (A) ORGANISM: PCR PRIMER

```
(vii) IMMEDIATE SOURCE:
                  (B) CLONE: CD40 3' PRIMER
   5
            (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:
       CCGTCGACGT CTAGAGCCGA TCCTGGGG
  10
       (2) INFORMATION FOR SEQ ID NO:7:
            (i) SEQUENCE CHARACTERISTICS:
                  (A) LENGTH: 40 base pairs
  15
                  (B) TYPE: nucleic acid
                  (C) STRANDEDNESS: single
                  (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: cDNA
 20
          (iii) HYPOTHETICAL: NO
           (vi) ORIGINAL SOURCE:
                 (A) ORGANISM: PCR PRIMER
 25
          (vii) IMMEDIATE SOURCE:
                 (B) CLONE: CD40 3' DOWNSTREAM PRIMER
 30
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:
      ACAAGATCTG GGCTCTACGT ACTCAGCCGA TCCTGGGGAC
      40
 35
      (2) INFORMATION FOR SEQ ID NO:8:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 5 amino acids
(B) TYPE: amino acid
40
                (D) TOPOLOGY: linear
          (ii) MOLECULE TYPE: peptide
         (iii) HYPOTHETICAL: NO
45
           (v) FRAGMENT TYPE: internal
          (vi) ORIGINAL SOURCE:
                (A) ORGANISM: PENTAPEPTIDE
50
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
          Tyr Val Gly Pro Arg
55
     (2) INFORMATION FOR SEQ ID NO:9:
```

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 43 base pairs(B) TYPE: nucleic acid

60

```
(C) STRANDEDNESS: single
                  (D) TOPOLOGY: linear
            (ii) MOLECULE TYPE: cDNA
   5
           (iii) HYPOTHETICAL: NO
            (vi) ORIGINAL SOURCE:
                  (A) ORGANISM: PCR PRIMER
  10
           (vii) IMMEDIATE SOURCE:
                  (B) CLONE: HUMAN IGG1/FC 5' PRIMER
  15
           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:
       TATTAATCAT TCAGTAGGGC CCAGATCTTG TGACAAAACT CAC
       43
  20
       (2) INFORMATION FOR SEQ ID NO:10:
            (i) SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 38 base pairs
                 (B) TYPE: nucleic acid
 25
                 (C) STRANDEDNESS: single
                 (D) TOPOLOGY: linear
           (ii) MOLECULE TYPE: cDNA
 30
          (iii) HYPOTHETICAL: NO
          (iv) ANTI-SENSE: NO
          (vi) ORIGINAL SOURCE:
 35
                (A) ORGANISM: PCR PRIMER
         (vii) IMMEDIATE SOURCE:
                (B) CLONE: HUMAN IGG1/FC 3' DOWNSTREAM PRIMER
40
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
      GCCAGCTTAA CTAGTTCATT TACCCGGAGA CAGGGAGA
      38
45
      (2) INFORMATION FOR SEQ ID NO:11:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 840 base pairs
50
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: cDNA
55
        (iii) HYPOTHETICAL: NO
         (iv) ANTI-SENSE: NO
60
         (vi) ORIGINAL SOURCE:
```

(A) ORGANISM: Homo sapiens

(vii)	IMME	DIATE	S	OURCE:
	(B)	CLONE		CD40-T

5 (ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 46..831

10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

TGCCACCTTC TCTGCCAGAA GATACCATTT CAACTTTAAC ACAGC ATG ATC GAA 54

Met Ile Glu

ACA TAC AAC CAA ACT TCT CCC CGA TCT GCG GCC ACT GGA CTG CCC ATC

Thr Tyr Asn Gln Thr Ser Pro Arg Ser Ala Ala Thr Gly Leu Pro Ile
5 10 15

AGC ATG AAA ATT TTT ATG TAT TTA CTT ACT GTT TTT CTT ATC ACC CAG $150\,$

25 Ser Met Lys Ile Phe Met Tyr Leu Leu Thr Val Phe Leu Ile Thr Gln 35

ATG ATT GGG TCA GCA CTT TTT GCT GTG TAT CTT CAT AGA AGG TTG GAC 198

30 Met Ile Gly Ser Ala Leu Phe Ala Val Tyr Leu His Arg Arg Leu Asp 40 45 50

AAG ATA GAA GAT GAA AGG AAT CTT CAT GAA GAT TTT GTA TTC ATG AAA 246

Lys Ile Glu Asp Glu Arg Asn Leu His Glu Asp Phe Val Phe Met Lys
55 60 65

ACG ATA CAG AGA TGC AAC ACA GGA GAA AGA TCC TTA TCC TTA CTG AAC 294

Thr Ile Gln Arg Cys Asn Thr Gly Glu Arg Ser Leu Ser Leu Leu Asn 70 75 80

TGT GAG GAG ATT AAA AGC CAG TTT GAA GGC TTT GTG AAG GAT ATA ATG 342

Cys Glu Glu Ile Lys Ser Gln Phe Glu Gly Phe Val Lys Asp Ile Met 85 90 95

TTA AAC AAA GAG GAG ACG AAG AAA GAA AAC AGC TTT GAA ATG CAA AAA 390

Leu Asn Lys Glu Glu Thr Lys Lys Glu Asn Ser Phe Glu Met Gln Lys 115

GGT GAT CAG AAT CCT CAA ATT GCG GCA CAT GTC ATA AGT GAG GCC AGC 438

Gly Asp Gln Asn Pro Gln Ile Ala Ala His Val Ile Ser Glu Ala Ser 120 125 130

AGT AAA ACA ACA TCT GTG TTA CAG TGG GCT GAA AAA GGA TAC TAC ACC

Ser Lys Thr Thr Ser Val Leu Gln Trp Ala Glu Lys Gly Tyr Tyr Thr 135

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	534	Į								•						C GTT
5		: Sei	150		. Leu	Val	Thr	155		Asn	ı GIŞ	r Lys	160		ı Thi	c Val
_	AAA 582															TCC
10	_	165		Gly	Leu	Tyr	Tyr 170	Ile	Tyr	Ala	Gln	Val 175		Phe	: Cys	Ser
10			GAA	GCT	TCG	AGT	CAA	GCT	CCA	TTT	ATA	GCC	AGC	CTC	TGC	CTA
15	Asn 180	Arg	Glu	Ala	Ser	Ser 185	Gln	Ala	Pro	Phe	Ile 190	Ala	Ser	Leu	Cys	Leu 195
	AAG 678	TCC	ccc	GGT	AGA	TTC	GAG	AGA	ATC	TTA	CTC	AGA	GCT	GCA	AAT	ACC
20	Lys	Ser	Pro	Gly	Arg 200	Phe	Glu	Arg	Ile	Leu 205	Leu	Arg	Ala	Ala	Asn 210	Thr
	726		TCC													
25	His	Ser	Ser	Ala 215	Lys	Pro	Cys	Gly	Gln 220	Gln	Ser	Ile	His	Leu 225	Gly	Gly
	GTA 774	TTT	GAA	TTG	CAA	CCA	GGT	GCT	TCG	GTG	TTT	GTC	AAT	GTG	ACT	GAT
30	Val	Phe	Glu 230	Leu	Gln	Pro	_	Ala 235	Ser	Val	Phe		Asn 240	Val	Thr	Asp
50	CCA 822	AGC	CAA	GTG .	AGC (CAT	GGC 2	ACT	GGC	TTC	ACG	TCC	TTT	GGC	TTA	CTC
35		Ser 245	Gln '	Val	Ser 1		Gly ' 250	Thr (Gly :	Phe		Ser 255	Phe	Gly	Leu	Leu
	AAA 840 Lys		TGAA	CAGT	GT CZ	Ā										
40	260	Беа														
	(2)	INFO	RMATI	ON I	FOR S	EQ I	D NC	:12:	:							
45		(:	i) SE	(A) (B)	CE C LENG TYPE TOPO	TH:	261 ino	amin acid	o ac	ids						
50			.) MO							TD N	n:12	•				
	Met I								er P				la A	la I	hr 0	Sly
55	Leu P	ro I		er M 20	et L	ys I.	le P		et T 25	yr L	eu L	eu T	hr V	al P 30	he I	eu
60	Ile T		ln Me 35	et I	le G	Ly Se		la Le 10	eu Pl	he A	la V		yr L 45	eu H	is A	ırg

	vrč	50		, гуде	s ite	: GIU	55		ı Arç	J Asn	ı Lev	His 60		ı Asr) Phe	e Val
5	Phe 65	e Met	Lys	Thr	lle	Gln 70	Arg	Cys	Asr	Thr	Gly 75		Arg	ser Ser	Leu	Ser 80
	Leu	Leu	Asn	Cys	Glu 85	Glu	Ile	Lys	Ser	Gln 90		Glu	Gly	Phe	• Val 95	
10	Asp	lle	Met	Leu 100	Asn	Lys	Glu	Glu	Thr 105		Lys	Glu	Asn	Ser 110		Glu
15	Met	Gln	Lys 115	Gly	Asp	Gln	Asn	Pro 120		Ile	Ala	Ala	His 125		Ile	Ser
	Glu	Ala 130	Ser	Ser	Lys	Thr	Thr 135	Ser	Val	Leu	Gln	Trp 140	Ala	Glu	Lys	Gly
20	Tyr 145	Tyr	Thr	Met	Ser	Asn 150	Asn	Leu	Val	Thr	Leu 155	Glu	Asn	Gly	Lys	Gln 160
	Leu	Thr	Val	Lys	Arg 165	Gln	Gly	Leu	Tyr	Tyr 170	Ile	Tyr	Ala	Gln	Val 175	Thr
25	Phe	Cys	Ser	Asn 180	Arg	Glu	Ala	Ser	Ser 185	Gln	Ala	Pro	Phe	Ile 190	Ala	Ser
30			195		Ser			200					205		_	
		210	,	,	Ser		215					220				
35	225					230					235					240
40					Ser (245	Gln	Val	Ser		Gly 250	Thr	Gly	Phe		Ser 255	Phe
40	Gly	Leu		Lys : 260	Leu											
45	(2)				FOR S											
		(i)	(A) (B)	LEI TYI	E CHA NGTH: PE: r	: 73	base	e pa:	irs						•	
50			(D)	TOE	RANDE	Y: 1	linea	ır								
					TYP			to m	nRNA							
55					ISE:								-			
60	(xi)	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	13:					

```
54
       TGGTGGCGGA GGGTCAGGCG GAGGTGGGTC CGGAGGCGGG GGTTCAAGTT CTGACAAGAT
       AGAAGATGAA AGG
  5
       73
       (2) INFORMATION FOR SEQ ID NO:14:
            (i) SEQUENCE CHARACTERISTICS:
 10
                 (A) LENGTH: 21 base pairs
                 (B) TYPE: nucleic acid
                 (C) STRANDEDNESS: single
                 (D) TOPOLOGY: linear
 15
           (ii) MOLECULE TYPE: cDNA
          (iii) HYPOTHETICAL: NO
          (iv) ANTI-SENSE: NO
 20
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:
25
      GGCCGCTCAG AGTTTGAGTA A
      (2) INFORMATION FOR SEQ ID NO:15:
30
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 1425 base pairs
                (B) TYPE: nucleic acid
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: linear
35
          (ii) MOLECULE TYPE: cDNA to mRNA
         (iii) HYPOTHETICAL: NO
40
         (iv) ANTI-SENSE: NO
        (vii) IMMEDIATE SOURCE:
               (B) CLONE: Human CD40-L/FC2 (soluble CD40-L)
45
         (ix) FEATURE:
               (A) NAME/KEY: CDS
               (B) LOCATION: 4..1422
         (ix) FEATURE:
```

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55

60

(A) NAME/KEY: mat_peptide (B) LOCATION: 79..1422

(ix) FEATURE:

(A) NAME/KEY: sig_peptide (B) LOCATION: 4..78

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

	T2 48	AT A	TG	TTC	CA!	r GI	T TC	T T	TT A	GA	TAT	ra 1	C T	T G	GA	AT	r cc	T C	CA	CTG
5		M -	et 1 25	Phe	Hi:	s Va	l Se		ne A 20	.rg	Туг	: Il	e Ph		1y 15	Ile	e Pr	o P	ro	Leu
3		C C	TT (ЭTT	CTC	G CT	G CC	T G1	C A	СT	AGC	rc	T GA	CT.	AC	AAA	GA	T G	АC	GAT
10	I1 -1	e L	eu t	/al	Let	ı Le	u Pr	o Va 5	al T	hr	Ser	: Se	r As	р Т 1	yr	Lys	As	p A	sp 5	Asp
10	GA 14	T A	AA A	AGA	TCI	TG'	T GA	C AA	A A	CT	CAC	AC.	A TG	C C	CA	CCG	TG	c c	CA	GCA
15	As	p Ly	rs A	rg	Ser 10	Cya	s As	р Ьу	s T	hr	His 15		r Cy	s Pi	ro	Pro	Cy.		ro	Ala
13	CC ¹	т G <i>‡</i> 2	AA C	TC	CTG	GGG	G GG2	A CC	G T	CA	GTC	TT	CT	C TI	rc	ccc	CC	A A	LΑ	ccc
20	Pr	o G1	u L	eu 25	Leu	Gl	/ Gly	, Pr	o Se	er 30	Val	Phe	e Le	u Ph	ıe	Pro 35	Pro	L.	75	Pro
20	AA(24(G GA	C A	CC	CTC	ATG	ATC	TC	c ce	G.	ACC	CCI	GA(G GI	c.	ACA	TGO	GI	'G	GTG
25	Lys	5 As 4	рТ 0	hr	Leu	Met	Ile	Se 4	r Ar 5	g	Thr	Pro	Glı		0	Thr	Cys	v a	1	Val
23	GT0	GA	C G	TG	AGC	CAC	GAA	GA	c cc	T (GAG	GTC	AAG	TT	C I	AAC	TGG	TA	С	GTG
30	Val 55	As	p Va	al	Ser	His	Glu 60	Asp	Pr	0 (Glu	Val	Lys 65		e 2	Asn	Trp	ту	r	Val 70
50	336	,					CAT													
35	Asp	Gl	y Vā	al (Glu	Val 75	His	Asr	Al.	a I	Lys	Thr 80	Lys	Pro	o A	rg	Glu	G1: 8:		Gln
	364						CGG													
40	Tyr	Ası	ı Se	er !	Thr 90	Tyr	Arg	Val	Va.	l S	er 95	Val	Leu	Thi	r V		Leu 100	His	3 (Gln
	432						AAG													
45	Asp	Trp	10	u <i>F</i> 5	lsn	Gly	Lys	Glu	Ту: 110	: L	ys (Cys	Lys	Val		er . 15	Asn	Lys	: A	la
	480						GAG													
50	ren	120	A1.	a P	'ro	Ile	Glu	Lys 125	Thr	: I.	le S	Ser	Lys	Ala 130		ys (Sly	Gln	P	ro
	CGA 528																			
55	Arg 135	GIU	PL	J	TH /	/aı	Tyr 140	Thr	Leu	P	ro P		Ser 145	Arg	As	sp (Slu	Leu		hr 50
	AAG 576																	• •		
60	Lys	11011	GII	. V	aı S 1	er . .55	ren .	rnr	cys	₽€		al : 60	Lys	Gly	Ph	e I		Pro 165	S	er

56

	GA 62		TC G	CC	GT	G GA	G TG	G G	AG	AGO	C AA	T G	GG	CAG	CC	G G	AG	AA	C AA	C	TAC
	As		le A	la	Va]		u Tr	p G	lu	Sei	As 17		ly	Gln	Pr	o G	lu	As:	_	n	Tyr
5	AA		CC A	.CG	CCI	cc	C GT	G C	TG	GAC	TC	C G2	AC	GGC	TC	C T	TC	TTC	C CT	С	TAC
	67 Ly			hr 85	Pro	Pro	o Va	l L	eu	Asp		r As	sp	Gly	Se		he 95	Phe	e Le	u '	Tyr
10	AG	C AA			ACC	GTO	GA	C A	AG			3 TC	G.	CAG	CAG			AAC	GT	c :	TTC
	72	0					. As														
15	mC:	20		~~	CERC	3 m/c)5	cca	CCI			m C 3	210		۰,	CCT		~ .	ncc
	768	3					CA:									_			_		
20	215	_			•••		220		·1	OL _J	رس	01	_	225	ريد	0.2	- <u>J</u>		02,	•	230
	816	5					AGI														
25	Gly	GL;	y G	ГĀ	Gly	Ser 235	Ser	: Se	r	Asp	Lys	11 24		3lu	Asp	Gl	u i	Arg	Asr. 245		eu
20	CAT 864		A GA	T '	TTT	GTA	TTC	AT	G 2	AAA	ACG	AT	A C	CAG	AGA	TG	C Z	AAC	ACA	G	GA
20	His	Glı	ı As		Phe 250	Val	Phe	Me	t I	Lys	Thr 255	Ile	e G	ln	Arg	Су		Asn 260	Thr	G	ly
30			A TC	c:	ΓΤΑ	TCC	TTA	CT	G P	AAC	TGT	GA	3 G	AG .	ATT	AA	A Z	AGC	CAG	T	TT
	912 Glu		s Se 26		Leu	Ser	Leu	Le		Asn 270	Cys	Glu	ı G	lu	Ile	Ly:		er	Gln	P	he
35	GAA	GGC			FTG	AAG	GAT	ATZ			TTA	AAC	: A.	AA (GAG			.CG	AAG	Αž	AA
	960 Glu	Gly	Ph					Ile	e M					ys (Glu						
40	CAA	280		~ m	mm	C	3.00	285			00m	~ 3 ***			290				3 mm	-	20
	GAA 1008 Glu	3																			
45	295						300			,,,,	J-y	110p		05						31	
	GCA 1056	;																			
50	Ala	His	Va]	L I.		Ser 315	Glu	Ala	S	er S		Lys 320	Th	ır I	hr	Ser	V		Leu 325	G1	n
50	TGG 1104		GAA	A	AA G	GA '	TAC	TAC	A	CC A	TG.	AGC	AA	C A	AC	TTG	G:	CA Z	ACC	CT	G
	Trp .		Glu		ys G 30	ly :	ryr	Tyr	Tì		let 135	Ser	As	n A	.sn	Leu	Va 34		lhr	Le	u
55	GAA 2	AAT	GGG	A.	AA C	AG (CTG .	ACC	GI	rt À	AA Z	AGA	CA	A G	GA (CTC	TA	T T	TAT	AT(С
	1152 Glu 1	Asn			7s G	ln 1	eu '	Thr			ys i	Arg	Gl	n G			ту	rr I	'yr	Ile	е
60			345						35	O					•	355					

57 TAT GCC CAA GTC ACC TTC TGT TCC AAT CGG GAA GCT TCG AGT CAA GCT 1200 Tyr Ala Gln Val Thr Phe Cys Ser Asn Arg Glu Ala Ser Ser Gln Ala 365 5 CCA TTT ATA GCC AGC CTC TGC CTA AAG TCC CCC GGT AGA TTC GAG AGA 1248 Pro Phe Ile Ala Ser Leu Cys Leu Lys Ser Pro Gly Arg Phe Glu Arg 10 ATC TTA CTC AGA GCT GCA AAT ACC CAC AGT TCC GCC AAA CCT TGC GGG 1296 Ile Leu Leu Arg Ala Ala Asn Thr His Ser Ser Ala Lys Pro Cys Gly 395 400 15 CAA CAA TCC ATT CAC TTG GGA GGA GTA TTT GAA TTG CAA CCA GGT GCT 1344 Gln Gln Ser Ile His Leu Gly Gly Val Phe Glu Leu Gln Pro Gly Ala 410 20 TCG GTG TTT GTC AAT GTG ACT GAT CCA AGC CAA GTG AGC CAT GGC ACT Ser Val Phe Val Asn Val Thr Asp Pro Ser Gln Val Ser His Gly Thr 25

GGC TTC ACG TCC TTT GGC TTA CTC AAA CTC TGA

Gly Phe Thr Ser Phe Gly Leu Leu Lys Leu 440 445

30

35

60

- (2) INFORMATION FOR SEQ ID NO:16:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 473 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- 40 (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

Met Phe His Val Ser Phe Arg Tyr Ile Phe Gly Ile Pro Pro Leu Ile
-25
-20
-15
-10

Leu Val Leu Pro Val Thr Ser Ser Asp Tyr Lys Asp Asp Asp Asp -5

Lys Arg Ser Cys Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro 10 15 20

Glu Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys 25 30 35

Asp Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val 40 45 50 55

Asp Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp 60 65 70

Gly Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr

75	. 80	85

) o	. 50	∽ ምክ	γ - Ψτσ	- A-	+ Val	Va	l Sei	r Vai	l Le	u Thi	r Val	L Le	ı Hi:	s Glı	n Asp
5	ASI	1 36	9		r ar	, ,		9:					10	0		•
J	Tr	10		n Gly	y Lys	s Glu	110		з Су:	s Ly	s Val	115		n Ly:	s Ala	a Leu
10	Pro 120		a Pr	o Ile	e Glı	125		r Ile	e Se	c Ly:	3 Ala 130		Gly	y Glr	n Pro	135
	Glu	ı Pro	Gl:	n Val	140		Let	ı Pro	Pro	Se:		, Asp	Glu	ı Leı	1 Thr 150	Lys
15	Asn	ı Glı	ı Val	155		Thr	Cys	Leu	Val 160		s Gly	Phe	туг	165		Asp
20	Ile	Ala	170		Trp	Glu	Ser	175		Glr	Pro	Glu	180		Tyr	Lys
20	Thr	Th:) Pro	Val	Leu	Asp 190		Asp	Gly	Ser	Phe 195		Leu	Tyr	Ser
25	Lys 200		Thr	Val	Asp	Lys 205	Ser	Arg	Trp	Gln	Gln 210		Asn	Val	Phe	Ser 215
	Cys	Ser	Val	. Met	His 220	Gly	Gly	Gly	Gly	Ser 225		Gly	Gly	Gly	Ser 230	Gly
30	Gly	Gly	Gly	Ser 235		Ser	Asp	Lys	Ile 240	Glu	Asp	Glu	Arg	Asn 245	Leu	His
35	Glu	Asp	Phe 250	Val	Phe	Met	Lys	Thr 255	Ile	Gln	Arg	Cys	Asn 260	Thr	Gly	Glu
55	Arg	Ser 265		Ser	Leu	Leu	Asn 270	Cys	Glu	Glu	Ile	Lys 275	Ser	Gln	Phe	Glu
40	Gly 280	Phe	Val	Lys	Asp	Ile 285	Met	Leu	Asn	Lys	Glu 290	Glu	Thr	Lys	Lys	Glu 295
	Asn	Ser	Phe	Glu	Met 300	Gln	Lys	Gly	Asp	Gln 305	Asn	Pro	Gln	Ile	Ala 310	Ala
45	His	Val	Ile	Ser 315	Glu	Ala	Ser	Ser	Lys 320	Thr	Thr	Ser	Val	Leu 325	Gln	Trp
50	Ala	Glu	Lys 330	Gly	Tyr	Tyr	Thr	Met 335	Ser	Asn	Asn	Leu	Val 340	Thr	Leu	Glu
50	Asn	Gly 345	Lys	Gln	Leu		Val 350	Lys	Arg	Gln	Gly	Leu 355	Tyr	Tyr	Ile	Tyr
55	Ala 360	Gln	Val	Thr	Phe	Cys 365	Ser	Asn	Arg	Glu	Ala 370	Ser	Ser	Gln	Ala	Pro 375
	Phe	Ile	Ala	Ser	Leu 380	Суз	Leu	Lys	Ser	Pro 385	Gly	Arg	Phe	Glu	Arg 390	Ile
60	Leu	Leu		Ala 395	Ala	Asn '	Thr		Ser 400	Ser	Ala	Lys	Pro	Cys 405	Gly	Gln

	Gln	Ser	Ile 410	His	Leu	Gly	Gly	Val 415	Phe	Glu	Leu	Gln	Pro 420	Gly	Ala	Ser	
5	Val	Phe 425	Val	Asn	Val	Thr	Asp 430	Pro	Ser	Gln	Val	Ser 435	His	Gly	Thr	Gly	
10	Phe 440	Thr	Ser	Phe	Gly	Leu 445	Leu	Lys	Leu								
	(2)	INFO	RMA	CION	FOR	SEQ	ID N	10:17	7:								
15		(i)		3) TY		: 33 amin	ami	no a	CS: acids	i.							
20		(ii)	MOL	ECUL	E TY	PE:	pept	ide									
		(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	ои о	:17:						
25		Arg 1	Met	Lys	Gln	Ile 5	Glu	Asp	Lys	Ile	Glu 10	Glu	Ile	Leu	Ser	Lys 15	Ile
		Tyr	His	Ile	Glu 20	Asn	Glu	Ile	Ala	Arg 25	Ile	Lys	Lys	Leu	Ile 30	Gly	Glu
30		Arg															

CLAIMS

What is claimed is:

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- 1. An isolated DNA sequence encoding a CD40-L polypeptide that binds to CD40, wherein the DNA sequence is selected from the group consisting of:
- (a) nucleotides 46 through 828, 184 through 828, or 193 through 762 of SEQ ID NO:11;
- 10 (b) DNA sequences which detectably hybridize to one of the sequences of (a) or their complementary strands under moderate stringency conditions; and
 - (c) DNA sequences which, due to degeneracy of the genetic code, encode a polypeptide encoded by any of the foregoing DNA sequences.
- An isolated DNA sequence according to claim 1 encoding a soluble CD40-L polypeptide.
 - 3. A recombinant expression vector comprising a DNA sequence according to claim 1.

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- 4. A recombinant expression vector comprising a DNA sequence according to claim 2.
- 5. A host cell transformed or transfected with an expression vector according25 to claim 3.
 - 6. A host cell transformed or transfected with an expression vector according to claim 4.
- 30 7. A process for preparing a CD40-L polypeptide, comprising culturing a host cell according to claim 5 under conditions promoting expression and recovering CD40-L polypeptide from the culture.
- A process for preparing a CD40-L polypeptide, comprising culturing a host
 cell according to claim 6 under conditions promoting expression and recovering CD40-L polypeptide from the culture.

9. A purified biologically active CD40-L polypeptide composition comprising a sequence of amino acids encoded by a nucleotide sequence according to claim 1.

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- 10. A purified biologically active CD40-L polypeptide composition according to claim 9, consisting essentially of human CD40-L.
 - 11. A purified biologically active CD40-L polypeptide composition according to claim 10, selected from the group consisting:
 - (a) the polypeptide defined by amino acids 1 through 261, inclusive, of the sequence set forth in SEQ ID NO:11;

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- (b) the polypeptide defined by amino acids 47 through 261, inclusive, of the sequence set forth in SEQ ID NO:11; and
- (c) a polypeptide defined by a sequence beginning with an amino acid in the sequence between amino acid 47 and amino acid 51, inclusive, through and including an amino acid in the sequence between amino acid 239 and amino acid 261, inclusive, of the sequence set forth in SEQ ID NO:11.
- 12. A purified biologically active CD40-L polypeptide composition according to claim 11, wherein the CD40-L is a soluble CD40-L.
- 13. A soluble CD40-L according to claim 12, wherein the soluble CD40-l comprises amino acids 51 through 261 of SEQ ID NO:11.
- 14. A soluble CD40-L according to claim 13 wherein the CD40-L is an
 25 oligomer comprising two or more CD40-L extracellular regions.
 - 15. A method of treating allergy, an allergic reaction, lupus, rheumatoid arthritis or graft versus host disease comprising administering an effective amount of a CD40 antagonist selected from the group consisting of a soluble CD40 protein, a CD40 fusion protein, a soluble monomeric CD40-L polypeptide, and combinations thereof.
 - 16. A method of augmenting a vaccine response with an adjuvant, wherein the adjuvant is selected from the group consisting of a membrane-bound CD40-L polypeptide, oligomeric soluble CD40-L polypeptides, and combinations thereof.
 - 17. A method for stimulating hybridoma cells to increase monoclonal antibody secretion, comprising administering an effective amount of a CD40 agonist selected

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from the group consisting of CD40-L, membrane-bound CD40-L and oligomeric CD40-L.

- 18 An antibody immunoreactive with CD40-L or a CD40-L immunogen.
- 19. An antibody according to claim 16 which is a monoclonal antibody.
- 20. An antisense or sense oligonucleotide that can inhibit transcription or translation of CD40-L, comprising a sequence of at least about 12 nucleotides
 10 corresponding to a DNA sequence according to claim 1 or its DNA or RNA complement.

1	TGCCACCTTCTCTGCCAGAAGATACCATTTCAACTTTAACACAGCATGATCGAAACATAC MetileGluThrTyr	60 5
6 <u>.1</u>	AACCAAACTTCTCCCCGATCTGCGGCCACTGGACTGCCCATCAGCATGAAAATTTTTATG	120
6	AsnGlnThrSerProArgSerAlaAlaThrGlyLeuProIleSerMetLysIlePheMet	25
121	TATTTACTTACTGTTTTTCTTATCACCCAGATGATTGGGTCAGCACTTTTTGCTGTGTAT	180
26	TyrLeuLeuThrValPheLeuIleThrGlnMetIleGlySerAlaLeuPheAlaValTyr	45
181	CTTCATAGAAGGTTGGACAAGATAGAAGATGAAAGGAATCTTCATGAAGATTTTGTATTC	240
45	LeuHisArgArgLeuAspLysIleGluAspGluArgAsnLeuHisGluAspFheValPhe	65
241	ATGAAAACGATACAGAGATGCAACACAGGAGAAAGATCCTTATCCTTACTGAACTGTGAG	300
65	MetLysThrileGlnArgCysAsnThrGlyGluArgSerLeuSerLeuLeuAsnCysGlu	85
301	GAGATTAAAAGCCAGTTTGAAGGCTTTGTGAAGGATATAATGTTAAACAAAGAGGAGACG	360
86	GluileLysSerGlnPheGluGlyPheValLysAspileMetLeuAsnLysGluGluThr	105
361	AAGAAAGAAAACAGCTTTGAAATGCAAAAAGGTGATCAGAATCCTCAAATTGCGGCACAT	420
106	LysLysGluAsnSerPheGluMetGlnLysGlyAspGlnAsnProGlnIleAlaAlaHis	125
421	GTCATAAGTGAGGCCAGCAGTAAAACAACATCTGTGTTACAGTGGGCTGAAAAAGGATAC	480
126	ValileSerGluAlaSerSerLysThrThrSerValLeuGlnTrpAlaAspLysGlyTyr	145
481	TACACCATGAGCAACAACTTGGTAACCCTGGAAAATGGGAAACAGCTGACCGTTAAAAGA	540
145	TyrThrMetSerAsnAsnLeuValThrLeuGluAsnGlyLysGlnLeuThrValLysArg	165
541	CAAGGACTCTATTATATCTATGCCCAAGTCACCTTCTGTTCCAATCGGGAAGCTTCGAGT	600
166	GlnGlyLeuTyrTyrIleTyrAlaGlnValThrPheCysSerAsnArgGluAlaSerSer	165
501	CAAGCTCCATTTATAGCCAGCCTCTGCCTAAAGTCCCCCGGTAGATTCGAGAGAATCTTA	660
L85	GlmAlaProFheIleAlaSerLeuCysLeuLysSerProGlyArgPheGluArgIleLeu	205
61	CTCAGAGCTGCARATACCCACAGTTCCGCCARACCTTGCGGGCAACAATCCATTCACTTG	720
206	LeuArgAlaAlaAsnThrHisSerSerAlaLysProCysGlyGlnGlnSerIleHisLeu	225
721	GGAGGAGTATTTGAATTGCAACCAGGTGCTTCGGTGTTTGTCAATGTGACTGATCCAAGC	780
226	GlyGlyValPheGluLeuGlnProGlyAlaSerValPheValAsnValThrAspProSer	245
81	CAAGTGAGCCATGGCACTGGCTTCACGTCCTTTGGCTTACTCAAACTCTGAACAGTGTCA	840
245	GlnValSerHisGlyThrGlyPheThrSerPheGlyLeuLeuLysLeu	261
341	CCTTGCAGGCTGTGGTGGAGCTGACGCTGGGAGTCTTCATAATACAGCACAGCGGTTAAGC	900
01	CCACCCCCTGTTAACTGCCTATTTATAACCCTAGGATCCTCCTTATGGAGAACTAETTAE	960

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100
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                                          OPSQVSHG
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3ASQVIHR
KTTSVLQWAEKGYYTMSN
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FIG. 4A

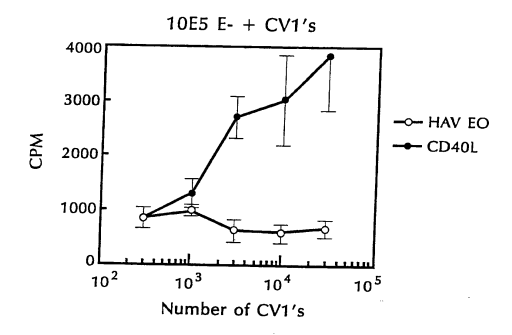
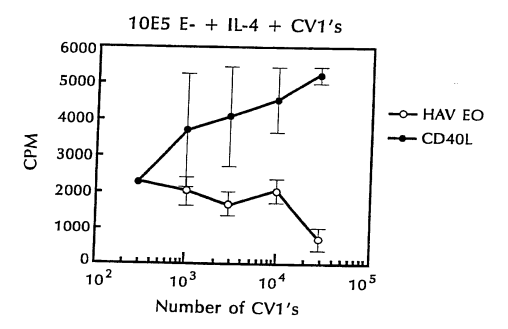


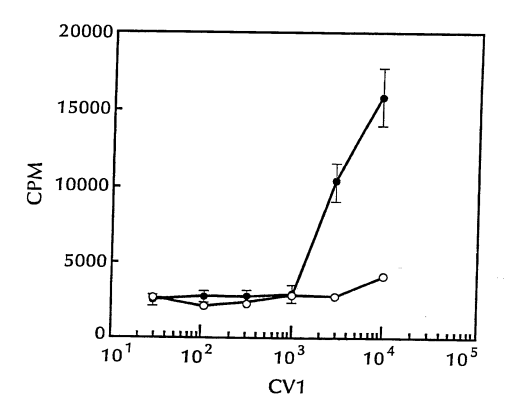
FIG. 4B



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FIG. 5

PB E + IL-4 + CV1 d7 Proliferation

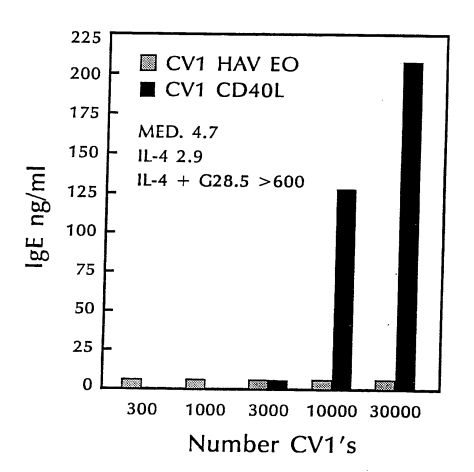


- → HAV **-** CD40L

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FIG. 6

S.CD23 in Day 6 Cultures S/N: 10E5 E-/Well, IMDM + IL-4



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FIG. 7

S.CD23 in Day 6 Cultures S/N: 10E5 E-/Well, IMDM + IL-4

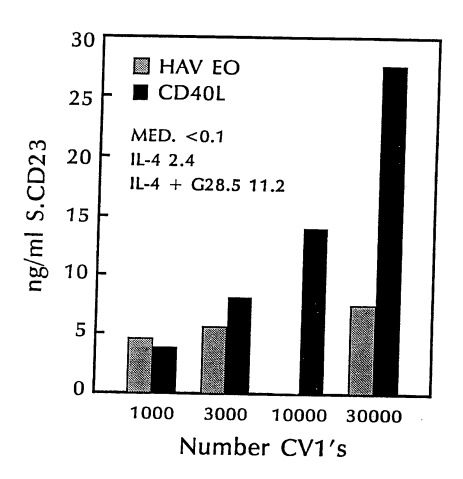
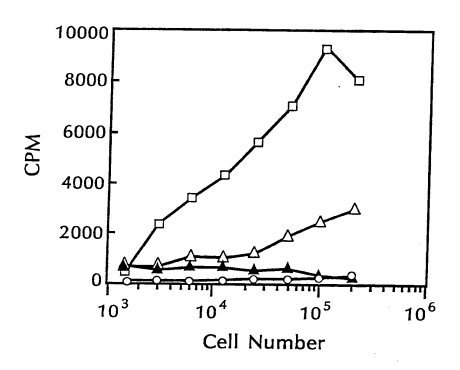


FIG. 8

Induction of B Cell Proliferation by CD40 Ligand Expressing CV-1 Cells (fixed)



---- CD40L

-△- 7A1

→ HAV EO

- 40.9

Induction of Anti-SRBC PFC by EL4 40.9 and 7A1 Th1 Cells (Fixed)

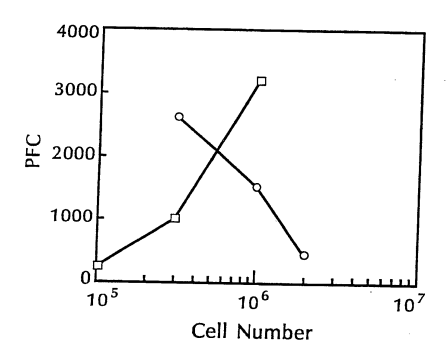
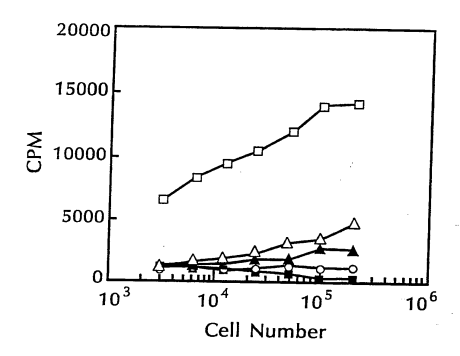


FIG. 10

Induction of B Cell Proliferation by CD40 Ligand Expressing CV-1 Cells



----- CD40L

-△- 7A1

--- HAV EO

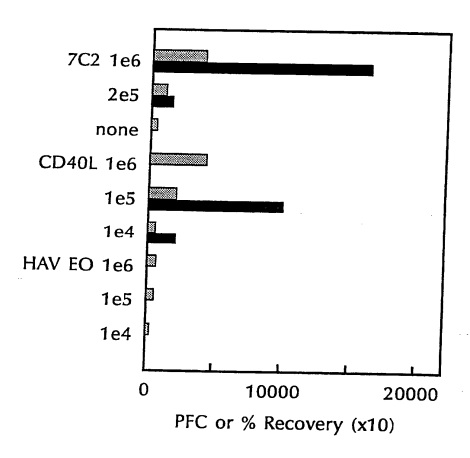
- 7A1 + CD40Fc

->- CD40Fc

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FIG. 11

Induction of Anti-SRBC PFC by CD40 Ligand Expressing CV-1 Cells (fixed)



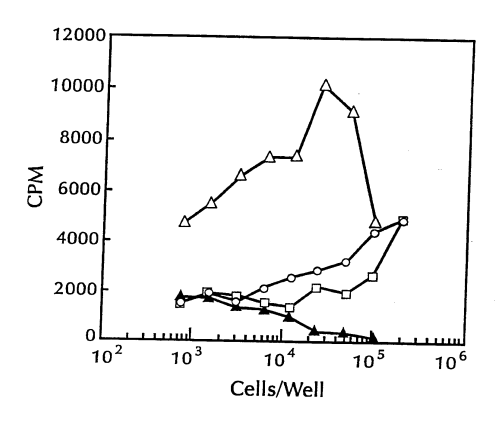
- % Recovery
- PFC

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FIG. 12

Induction of Murine B Cell Proliferation by CD40 Ligand Expressing CV-1 Cels (fixed)



-∆- CD40L

→ HAV EO

-□- 7A1 11/6

13/14 FIG. 13A

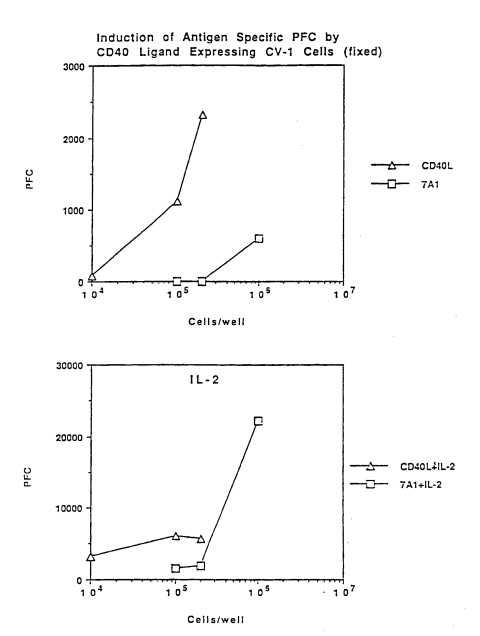


FIG. 13B

FIG. 14A

Day 7 Proliferation of T-depleted PBM

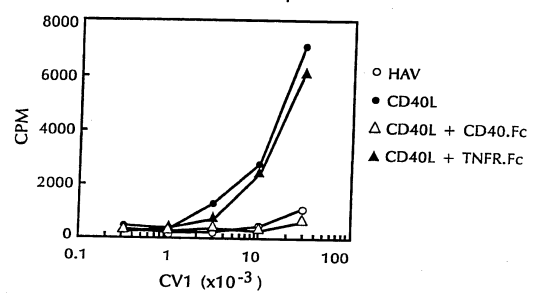
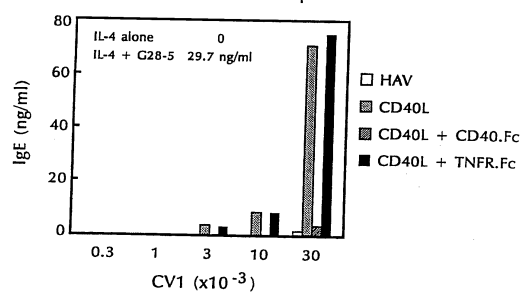


FIG. 14B

Day 10 IgE Secretion from T-depleted PBM



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INTERNATIONAL SEARCH REPORT

International application No. PCT/US92/08990

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A. CI IPC(5)	ASSIFICATION OF SUBJECT MATTER :Please See Extra Sheet.								
US CL									
According	g to International Patent Classification (IPC) or to	both national classification and IPC							
	ELDS SEARCHED								
Minimum documentation searched (classification system followed by classification symbols)									
U.S. : 536/27; 530/388.22, 388.23, 389.2, 351; 435/70.3, 69.1, 69.5, 240.1, 252.3									
		0.5, 09.1, 09.5, 240.1, 252.3							
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APS, DIALOG									
search terms: CD40, ligand, antibodies, recombinant									
C. DOC	CHAPTE CONCIDENCE TO BE DELEVISION								
	CUMENTS CONSIDERED TO BE RELEVAN		T						
Category*	Citation of document, with indication, when	re appropriate, of the relevant passages	Relevant to claim No.						
	Nature, Volume 350, Issued 04 April 1991, M	.L. Riordan et al., "Oligonucleotide-based	20						
	therapeutics", pages 442-443, entire document.								
7	The EMBO Journal, Volume 8(5), issued 1989	9 I Stamenkovic et al. "A D lumphouse							
	activation molecule related to the nerve growth	factor recentor and induced by cytokings in	1-14, 17-20						
carcinomas", pages 1403-1410, entire document, especially page 1408 2nd column									
	paragraphs 2-3.								
ĺ	Proceedings of the National Academy of Scien	nce, Vol. 84, issued December 1987 A	1-6						
	Aruno and B. Seed, "Molecular cloning of a C	D28 cDNA by a high-efficiency COS call	1-0						
l	expression system", pages 8573-8577, entire doc	cument, see especially page 8573, §2.							
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Further	r documents are listed in the continuation of Box	C. See patent family annex.							
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US92/08990

									
A. CLASSIFICATION OF SUBJECT MATTER: IPC (5):	·								
C07H 21/00: A61K 35/14; C07K 3/00, 7/00, 13/00; C12P 21/02, 21/06; C12N 15/00									
A. CLASSIFICATION OF SUBJECT MATTER: US CL:									
536/27; 530/388.22, 388.23, 389.2, 351; 435/70.3, 69.1, 69.5, 240.1, 252.3									
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